Congratulations!

With debritom®, you have acquired a high quality and innovative micro water jet device that sets new standards for wound debridement. debritom® has an electronic control system with optical status indications. Most components that will be getting in contact with rinsing fluids are single use, which have to be disposed of after use according to your internal guidelines. The size enables high flexibility and mobile use, and the extreme quietness of operation allows the user to concentrate on wound and patient for an efficient and safe debridement. The comprehensive range of accessories offers the medical professional a toolbox for their individual needs.

1 Warnings and safety instructions

WARNINGS
Indicates a potentially hazardous situation which, if not avoided, could result in death or serious injury.

CAUTION
Indicates a potentially hazardous situation which, if not avoided, could result in minor or moderate injury.

SAFETY INSTRUCTION
Indicating useful information about the safe use of the device.

debrition® is approved exclusively for the use as described in these instructions for use. Medaxis can only guarantee the safe functioning of the system when debritom® is used in combination with the original debritom® accessories (pump, handpiece, connection tube, power cord; see chapter 17 – Accessories).

debrition® is EMC-tested in conformity with the requirements of IEC 60601-1-2:2014 and can be used in the vicinity of other EMC-tested devices that fulfill the requirements as outlined in the IEC 60601-1-2 standard. Untested HF sources, radio networks or the like can impair the function of the device and should not be operated in the vicinity of debritom®.

Please read and pay attention to the warning and safety instructions before operation. These instructions for use must be kept with the device for later reference.

Please note that these Instructions for use are a general guide for the use of the product. Medical matters must be addressed by a physician.

Medaxis considers itself only responsible for the effect on BASIC SAFETY, reliability and performance of debritom® if it is used in accordance with the Instructions for use.

Subject to change.

WARNING
– This manual must be read prior to the use of debritom®
– Please read and observe these warning and safety instructions before operation. These instructions for use must be kept with the device for later reference.
– The use of the device should only be carried out by medical, qualified personnel.
– Before cleaning the device you must disconnect the power supply.
– debritom® was validated in combination with the accessories listed in chapter 17. For a correct and safe operation use debritom® with these accessories only.
– Do not use debritom® in MR environment.
– The device is not suitable for use in a hazardous explosive environment.
– The use of debritom® for any other indication than intended is neither desired nor allowed.
– Wear gloves, protective googles and surgical masks for all operations.
– To avoid the risk of electric shock, this equipment must be connected to a power outlet with protective earth only.
– The device shall not be serviced or maintained while in use with a patient.
– Immediately disconnect the device from the supply mains and stop using it, if it is damaged.
– Modification of the device or combination with other devices is not allowed.
– Treating infants, make sure a second medical professional is holding the body to be treated securely in order to prevent sudden movement.
– Adults must be asked to keep the body to be treated securely in place.
– Not for use at home or in vehicles.

CAUTION
– The intensity level to be set must be determined by a physician according to the patient’s wound.
– Before connecting debritom® to the power supply, please verify that the supply voltage corresponds to that given on the device specification plate.
– Incorrect use of debritom® can cause pain and injury to the patient.
– Do not use sterile accessories when the sterile packaging is damaged.
– Non sterile and reusable accessories must be cleaned and disinfected according to chapter 10, General cleaning guidelines.
– US Federal law restricts this device to be sold by or on the order of a physician.
SAFETY INSTRUCTION

- debritom® is a medical device that requires specific safety measures in regard to EMC. It must be installed and put into operation in accordance with the attached EMC information in chapter 16, 'Technical documentation'.
- Portable and mobile RF communications equipment can affect medical devices.
- In each of the following cases, debritom® must not be used and it must be repaired by customer service:
  - If the power cord or the plug are damaged
  - If the device is not functioning according to routine check
  - If the device is damaged
  - If the device shows clear safety defects.
  - If a specific error reoccurs several times.
- debritom® has no user serviceable parts inside (open device with cover removed). For safety reasons, it is required that debritom® is repaired throughout its service life strictly and exclusively by Medaxis authorised service centres.
- Keep the power supply cord away from hot surfaces.
- The mains plug and the on/standby button must not come into contact with moisture. Never pull the mains plug out of the fixed mains socket by pulling on the power supply cord.
- Separation from the mains is only assured through the disconnection of the mains adapter and fixed socket connection.
- Never use the device at high room temperatures.
- Never place or immerse debritom® in water or other liquids.
- When using single use, sterile products, please note that they are not intended to be reprocessed. Reuse could result in loss of mechanical and chemical properties and may lead to restriction in terms of biocompatibility.
- Contact your local Medaxis customer service representative for assistance with product operations.
- The device can be switched off at any time by pulling out the mains plug.
- Do not operate the device when tired.

These instructions for use must be kept for later reference!

2 Power supply

debritom® is a mains-powered device. Before you plug in the device, please check that your local power supply is the same as the voltage given on the specification plate.

⚠️ WARNING
- To avoid the risk of electric shock, this equipment must be connected to a power outlet with protective earth only.
- Do not position the debritom® in away that it will be difficult to disconnect the power supply.

Plugging in the device
Take the power supply cord and plug in first the appliance inlet port at the back of the device.
3 Description

Introduction
debritom+ is a high precision micro water jet device. The compact system produces a fine waterjet to clean acute and chronic wounds in a precise and tissue-preserving manner. This highly concentrated and versatile micro water jet has multiple settings, for all water jet applications such as the gentle removal of foreign bodies via washing, irrigation, jet lavage and pulse lavage, for the removal of biofilm, necrotic and fibrotic tissue, and to generate micro bleeding tissue.

Using a sterile washing or debridement solution, a hand piece and pressure setting on the device are chosen to deliver the micro water jet therapy of choice. The gentleness of the wiping motions, the angle and the distance to the wound allow further refinement of the degree of wound irrigation or debridement applied. The practitioner, patient and surrounding area are protected from contamination by a protective cover.

The micro water jet therapy continues until the desired degree of irrigation or debridement is achieved. Upon completion of the micro water jet therapy, the gentle removal of foreign bodies, biofilm, necrotic and fibrotic tissue, or the creation of bleeding tissue, leaves the wound optimally prepared for healing.

debritom+ is intended for use on patients in appropriate care settings by a medical professional. (not for use in homecare environment nor in vehicles)

Intended purpose
debritom+ is intended for jet lavage, cleaning, irrigating, and debriding wounds and other diseases of the skin through the use of micro water jet technology.

Indications
Chronic, badly healing and stalled wounds which, in the clinical judgement of the responsible physician, require an irrigation or debridement procedure.
Acute wounds, which in the clinical judgement of the responsible physician, require an irrigation or debridement procedure.
Other diseases of the skin, which in the clinical judgement of the responsible physician, require an irrigation or debridement procedure.

Typical applications for use of debritom+
Infected, necrotic, ischemic, contaminated, or other poorly healing chronic wounds that, by example, may originate from the following diseases:
– Ulcers of various origin (venous, arterial, mixed)
– Ischemia
– Bedsores
– Diabetic foot syndrome
– Abscesses and fistulas
Acute wounds that, by example, may originate from the following:
– Accidents
– Burns
– Surgery
Removal of biofilm, of foreign bodies from wounds, eg. sand, textiles, fibers, metallic or wooden splinters, creation of microbleeding.

Contraindications
– Malignant tumors
– Open injuries to blood vessels, and unprotected blood vessels
– Eyes, ears, nose
– Delicate vessels and structures, such as neurovascular structures
– Complex and/or highly contaminated wounds
– Patients with HIV, Hepatitis C, and other contagious diseases

A warning with regard to the use of the device for the following conditions, since they are associated with higher risk. If treatment is indicated for the following conditions, they should only be performed under medical guidance and supervision:
– Patients with increased tendency to hemorrhage and arterial ulcers
– Infected wounds

Side-effects
There could be undesirable side-effects such as excessive bleeding or pain. In such cases it’s the physician’s responsibility to decide whether the debridement procedure shall be continued and under what circumstances (analgesic, intensity level decrease), or if the treatment shall be stopped.

Intended user
The debritom+ micro water jet therapy is designed for use by qualified health professionals.

Important note
Compliance with the proper irrigation and debridement procedures and techniques remains the responsibility of the physician. Each physician must evaluate the appropriateness of the treatment based upon their own knowledge and experience.
4 Overview debritom® system

debrtom®, REF 1000.0005
debritom® for recurrent use, for multiple patients

debritom® pump, REF 2000.020x
Pump versions depending on duration of use, either replace after application or after one day.

- Connector for connecting tube – rinsing fluid
- Luer-connector to handpiece
- Pump / drive unit interface to debritom®

Pole for rinsing fluid
Knurled head screw for pole
Display / User interface
Pump insert
Power connector / fuse
Specification plate and connection to trolley

- debritom® Pump
  REF 2000.0200
  single use, sterile

- debritom® Pump, one day
  REF 2000.0201
  one day use, multiple patients, sterile

With protective cap

debritom® contains items according to chapter 6
**debritom+ handpiece, REF 2000.000x**
Single use, sterile, various jet shapes, details see chapter 7 'Optimal working parameters'

**debritom+ connecting tube, REF 2000.0301**
Sterile, details see chapter 8 'Replace rinsing fluid'

**debritom+ foot on/off switch, REF 2000.5020**
For recurrent use, for multiple patients

**debritom+ power cord, REF 2000.5008**
Optional accessories

DebriClip    REF 2000.4001

Aeroguard "Jellyfish"    REF 2000.2000

Aeroguard "Nebbia"    REF 2000.2003

Connection port for handpiece

Connection port for suction device

Connection port for handpiece

Connection port for handpiece

Connection port for handpiece

Connection port for handpiece

Connection port for handpiece

Connection port for handpiece

Connection port for handpiece

Trolley    REF 1000.0500

5 Additional information

Intensity levels/running time

The handpiece and the pump are equipped with an RFID tag. These tags set according to the connected handpiece, with its corresponding water jet, a preset intensity level at the console. By connecting the handpiece, the intensity will be set at a standard level per handpiece. To increase the water jet’s intensity press ‘+’, to decrease the water jet intensity press ‘-’. See corresponding handpiece Instructions for intensity level and working distance.

Additionally, the RFID tags ensure a maximal running time of the pump and handpiece, in order to prevent wear and blockage of the system.
6 Installation

Check initial delivery
Check the delivery package of debritom+ for completeness and general condition.

**debritom+ basic unit**
- REF: 1000.0000
- with type label and transportation safety bolt

**Pole for rinsing fluid**
- REF: 8000.0001

**Knurled head screw for pole**
- REF: 8000.0000

**debritom+ power cord (US)**
- REF: 2000.5008

**debritom+ foot on/off switch**
- REF: 2000.5020

**debritom+ instruction for use**
- REF: 9000.5504

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Initial start up

Activate the foot on/off switch and check whether the green indicator light flashes.

Please note - Your foot on/off switch that comes with your device is already paired.
7 Preparation for use/operating instructions

**WARNING**
- The use of the device should only be carried out by medical qualified personnel.
- Wear gloves, protective goggles and surgical masks for all operations.
- Adapt intensity levels in the case of pain-sensitive patients.

**CAUTIONS**
- Do not use sterile accessories when the sterile packaging is damaged.
- Sterile products should be opened just before use.
- Non sterile and reusable accessories must be cleaned and disinfected according to chapter 10, 'General Cleaning Guidelines'.

**Frequently used functions**
- Insert pump
- Insert spike of connection tube into rinsing fluid bottle or bag
- Connect rinsing fluid with pump
- Connect handpiece to pump
- Open clamp and filter port (connection tube)
- Debridement of wound with micro water jet (jet on/off with foot pedal / intensity level / distance and angle of micro water jet to wound)
- Remove handpiece
- Remove pump

**Checks before use**

Check the debritom+ system before use for damage of the power cord or plug, obvious device damage or safety defects and proper functioning of the device.

- Check that the blue seal for the drive unit is correctly placed.
- Check all accessories prior to use:
  - handpiece, pump and connecting tube for cracks, brittle spots or other damages - replace if necessary.
- Check whether the blue LED lights up as the foot on/off switch will be activated. If needed replace batteries as shown in chapter 11 – Battery type 2 x LR03 / AAA

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**Connect debritom+ to mains**

**Turn on debritom+**

**General Button Operation**

debritom+ is a mains-powered device. Before you plug in the device, please check that your local power supply is the same as the voltage given on the specification plate.

- To avoid the risk of electric shock, this equipment must be connected to a power outlet with protective earth only.
- Do not position the debritom+ in away that it will be difficult to disconnect the power supply.
- Press any button for a short period of time to turn on the device or to adjust/change intensity settings.
- To switch off the device, press the on/standby button for 3 seconds.
- The on/standby button will also be used to acknowledge any type of error, to recognize as the button lights up red in the event of error messages. (see chapter 9)

**Note:** The intensity level of the micro water jet will be chosen automatically depending on type of handpiece (pre-programmed)

**debritom+ Display**

- Display for intensity levels and information
- Intensity level +
- Intensity level -
- on/standby button
**Connecting foot on/off switch with debritom⁺**

1. Press [ ] to turn on debritom⁺. The device immediately goes into the ‘Connecting mode’ for the foot on/off switch.

2. Connect the foot on/off switch with debritom⁺ by pressing the foot on/off switch once. The symbol ‘foot on/off switch connected’ is displayed and debritom⁺ is in standby mode.

If a connection to the foot on/off switch can not be established or a new foot on/off switch is used, this must be paired with the device. (Chapter 11 ‘Pairing foot on/off switch with debritom⁺’)

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**Insert debritom⁺ pump**

1a debritom⁺ pump: Open sterile blister pack and remove the pump.

1b debritom⁺ pump, one day: Open sterile blister pack and remove the pump. Keep the protective cap still in the blister pack. This will be used later as described under ‘Steps using debritom⁺ pump, one day REF 2000.0201’

2. Align triangle on pump with triangle on blue ring and insert pump horizontal as shown until it stops.

3. Turn pump clockwise until second triangle is aligned with the triangle on blue seal of drive unit. You will feel a ‘click’ as the pump reaches its final run position.

⚠️ **CAUTION**

When the pump is not installed correctly, a leakage of the treatment fluid could happen. In this case do not use the device any more and disconnect it from mains.
**Rinsing fluid**

The user does decide what type of rinsing fluid should be used to debride. These are generally saline solutions, or fluids containing polyhexanide. The fluid pole’s versatile hanger part is designed to attach the most common fluid bottles that are equipped with a shackle, fluid volumes up to 1000 ml. In chapter 17, you will find a list with rinsing fluids that are tested for compatibility with debritom+.

**NOTE:** Within 12 minutes of fulltime use, set on high intensity levels and no micro water jet interruption, you will approximately need 1000 ml of rinsing fluid.

**Attach the connecting tube**

1. Open sterile pouch.  
2. Make sure the clamp is in closed position.
3. Push the spike into the rinsing fluid.  
4. Open the venting cover.

**Connect the connecting tube with pump**

**Attach the debritom+ handpiece**

**WARNING**

Use the handpiece solely in combination with debritom+ pump and do not re-use (single use).

1. Open sterile pouch. Recommendation: due to hygienic reason, keep the handpiece within the pouch until start of treatment.
2. Attach the luer connection to the pump. Make sure the connection is tighten securely.
3. Open the clamp before start of treatment.
Placement of connected handpiece

The opening in the fluid pole serves as short time parking position for the handpiece during treatment interruptions. Insert the handpiece as shown.

**WARNING**
After each patient, the fluid pole, especially the opening for the handpiece needs to be disinfected thoroughly.

Further optional accessories

- Trolley see chapter 17.

- Aeroguard «Jellyfish» separate IFU 9000.5502.

- Aeroguard «Nebbia» separate IFU 9000.5506.

- DebriClip see chapter 27.
Wound treatment with debritom®

How it works
As you activate the foot on/off switch, the device starts running, resp. starts pumping rinsing fluid through the handpiece onto the patient’s wound.

The pressure to generate the micro water jet is being created inside the pump. The rinsing fluid will be pressed through the integrated nozzle within the handpiece, which creates the micro water jet. The regulation of the micro water jet’s intensity level is managed by the motor’s revolution.

Should the user realize, that the intensity level inadvertently increases, without any handling changes at the device, the foot on/off switch should not be pressed anymore, and the handpiece needs to be brought into a safe position, away from patient and user. The jet can also be interrupted by pressing any button on the debritom® device. Then debritom® needs to be switched off by pressing the on/standby button for 3 seconds. Should the user be unable to turn the device off, either the water supply or the electrical power needs to be disconnected.

All fluid conveying parts (handpiece, pump, connecting tube) are sterile and need to be disposed according to facility guidelines. This prevents cross contamination.

⚠️ Wound treatment steps

NOTE: Always wear protective gear according to internal guidelines during the treatment. Recommended are gloves, eye wear and surgical masks.

1. For chronic and algesic wounds it is recommendable to apply a local anesthetic. (Please follow the instruction of the manufacturer)

2. The patient needs to be in a comfortable position.

3. Put an absorbent cover underneath the wound location and make sure, that the rinsing fluid can easily drain off from the wound area.

4. Visually assess the micro water jet (correct shape: standard or flat, wide or narrow) as well as the jet’s intensity on your palm – distance approx. 20cm. If ok, move on to 5. If not ok, replace the handpiece and repeat 4.

5. For a patient that undergoes his first debritom® treatment, demonstrate the micro water jet also on his palm with a minimum distance of 8” / 20cm. This will increase his confidence into the new debridement treatment.

6. Start the treatment with either the pre-set intensity level for the chosen handpiece, or set your desired intensity level manually.

7. Press the foot on/off switch and start the procedure with the micro water jet tangentially to the wound, nozzle distance at working distance to the wound as prescribed below. Guide the jet starting at the wound edge, and move the jet slightly back and forth.

8. The optimal working angle lies between 3 and 45 degrees to the wound surface.

Optimal working parameters

Intensity levels at START of each handpiece : Level 4 (Level 1-5 / weak-strong)

debritom® handpiece standard jet
Working distance recommended 6” / 15-20cm

(debritom® handpiece flat jet narrow
Working distance recommended 1.5-4” / 4-10cm

(debritom® handpiece flat jet wide
Working distance recommended 1.2-3” / 3-8cm

NOTE flat jet: The flat jet should be applied like a spatula to the wound. (FIG)
Aerosol containment during wound treatment (optional accessories)

For further protection regarding aerosols, Medaxis recommends to use the AeroGuard Jellyfish as per separate instruction for use. Unfold the AeroGuard and use according to the instructions.

AeroGuard «Nebbia» is best suited for lower legs and arms. Do not use the product for any wounds on a head! This product protects user and patient from aerosols during mechanical wound cleansing/debridement.

The DebriClip is an additional attachment for debritom+ handpieces and is used to reduce aerosols and to collect rinsing fluid with debris during mechanical wound cleansing. (Detailed description p. 27)

Suitable for use with flat jet (wide and narrow).

for use with:
2000.0004
2000.0005

Wound treatment with DebriClip (aerosol insulation with mechanical wound cleansing)

Basis: debritom+ ready for use

Remove DebriClip from packaging
Open sterile package
Snap the handpiece into the opening provided

The jet can be aligned by turning the handpiece
Remove the DebriClip from the packaging with the engaged handpiece
Apply the DebriClip compact to the wound and debride according to IFU 9000.5503.

Additional function

4.1 Suction
Remove plug
Select and connect suitable suction hose
Start the suction system (select and connect a suitable suction system with a free flow rate of 10l/min to 30l/min)

4.2 Mechanical cleaning
Clean the wound and wound edges mechanically with the sponge

⚠️ CAUTION!
Not to be used with the standard jet REF 2000.0003, since the standard jet delivers a stronger and more intense water jet. If used in combination with DebriClip, the jet would be too close to the wound, much closer than the recommended minimum distance according to chapter «Optimal working parameter» (page 27).
Adjustment of water jet intensity

Adjust the Intensity level during the treatment
- Increase the distance between handpiece and wound => intensity level will decrease
- Decrease the distance between handpiece and wound => intensity level will increase

Adjust the intensity level at the device itself

⚠️ CAUTION
Before adjusting the intensity level, the treatment MUST be temporarily stopped. Continue with the treatment according to chapter 7 after adjusting the intensity level with +/-.

💡 SAFETY INSTRUCTION
Use a sterile gauze in order to absorb rinsing fluid.

To increase the water jet’s intensity
=> press +

To decrease the water jet intensity
=> press -

⚠️ CAUTION
- Switching off the device is possible at any time by pulling out the power plug.

After completion of the debridement procedure
As soon as the debridement procedure is completed, proceed with the following steps:

1. Press the ( for 3 seconds to set the debritom⁺ into standby mode and clamp the connecting tube.

2. Take care of your patient according to your facility’s protocol.

Steps after using debritom⁺ pump, single use REF 2000.0200

3a. Remove all single patient use components in following order:
- Remove the spike of the venting tube.
- Rotate the single patient use pump by 30 degrees counter clockwise and remove from the device. (There is no need to disconnect handpiece and connecting tube from the pump for disposal)
4a. Dispose of the complete unit (pump, handpiece and connecting tube) according to your facility guidelines. Follow the hospital's internal guidelines regarding period of use of the rinsing fluid.

4b. Use the protective cap to close the luer connection at the pump. (The cap can be found in the pump’s blister package.)

5a. If you are not planning on using debritom® soon again, disconnect the device from mains. (Pull power cord)

5b. Before attaching the next handpiece, remove the protective cap and store it in a clean place. (Opening downwards)

6. Unclamp connecting tube and start the treatment as described under “Wound treatment with debritom®”

Repeat steps 3b – 5b until your last treatment of the day. Then proceed with steps 3a – 5a.

⚠️ CAUTION
– Cover the pump’s luer connection with the cap immediately after removing the handpiece.

8 Replace rinsing fluid

If you should have to replace the rinsing fluid during a treatment, perform the following steps. Always make sure that the handpiece is being stored within a sterile setting and in a non critical position for patient and user.

1. Press 🔄 for 3 seconds to set the debritom® into standby mode.

2. Prepare the new rinsing fluid bottle in order to keep the time to switching over as short as possible.
3. Remove the spike of the connecting tube from the empty bottle and insert into the new bottle immediately. Make sure that the spike is not solid or contaminated. Unhook the empty bottle and attach the new one.

4. Press \(\text{ \text{on/standby button}}\) to turn on debritom\(^+\), and continue the treatment.

If the foot on/off switch should not be connected anymore, follow the instructions ‘connecting foot on/off switch’ in chapter 7.

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**CAUTION**

- Immediately connect the spike of the connecting tube to the new rinsing fluid bottle.

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9 Error messages

**NOTE:** The on/standby button lights up red as an error message occurs. This button can be used to acknowledge the fault after rectification.

- **Battery foot on/off switch almost empty**
  Troubleshooting: Replace batteries (Battery type 2 x LR03 / AAA, chapter 11)

- **Pump in use for too long time period**
  Troubleshooting: Replace pump

- **Handpiece in use for too long or trying to use with different single use pump**
  Troubleshooting: Replace handpiece

- **Handpiece missing or handpiece not recognized**
  Troubleshooting: Replace handpiece. If the error cannot be resolved, contact the Medaxis service department immediately.

- **Nozzle blocked or motor overheated**
  Troubleshooting: Change handpiece and pump if necessary. If the error cannot be resolved, contact the Medaxis service department immediately.

- **Pump not inserted correctly**
  Troubleshooting: Replace pump and reinsert it correctly, or replace it.

- **General system error**
  Troubleshooting: Please contact the Medaxis service department immediately.

- **No foot on/off switch detected**
  Troubleshooting: Press the foot on/off switch. If the foot on/off switch cannot be detected, replace the batteries (see chapter 11) and operate the foot pedal. If the foot on/off switch is still not detected, repeat the pairing procedure (see chapter 11)

- **Foot on/off switch time-out**
  The symbol appears if the foot on/off switch is pressed too slowly or the foot on/off switch is not pressed all the way down.
  Troubleshooting: Release the foot on/off switch completely to acknowledge the error. Press the foot pedal all the way down to start the water jet. If the fault cannot be rectified, contact the Medaxis service department immediately.
10 General cleaning guidelines

WARNING

After each use, the parts that have been in contact with aspirated secretions are to be cleaned and disinfected or disposed.

Before cleaning the device, pull the mains plug out of the fixed mains socket.

Safety instructions

- Never place electrical devices into water or other liquids.
- Do not spray or pour liquid directly onto debritom+.
- debritom+ product cannot be sterilized.
- Rinsing the debritom+ product range in a washing machine is not permitted.
- Immersion disinfection, thermal disinfection and ultrasound cleaning for the debritom+ product range are not permitted.
- Higher temperature than 45°C, 113°F can cause protein coagulation which can lead to problems further in the process.
- Clean all surfaces immediately after use to avoid residues from drying and to prevent growth of microorganisms.
- Do not use cleaning agent/detergent based on phenol.
- Do not use steel brushes or steel wool for cleaning.
- Store medical products dry and dust free.

Disposables

This Symbol indicates a single used product. This product is not intended to be reprocessed. Reuse could result in loss of mechanical and chemical properties and may lead to restrictions in terms of biocompatibility.

Exception – debritom+ pump, one day

The debritom+ pump, one day, is intended to be used for multiple patients within one day. The sterile package for the pump contains a luer cap, which is intended to cover the luer connection for the handpiece between switching patients.

After one day, the debritom+ pump, one day, has to be disposed of and is not intended to be reprocessed.

Reusable parts – debritom+, foot on/off switch

1. Thorough cleaning

Thorough cleaning can either be done at the point of use with sterile cold water only or in dedicated health care facilities with the additional use of enzymatic detergents according the manufacturer’s instructions for use.

CAUTION

Make sure that the wiping towel is not too wet, so that it does not drip onto the device.

1.1 Disconnect the power plug from the power socket.
1.2 Disinfect your hands and put on disposable gloves and proper protective gear.
1.3 Separate all individual parts. Dispose single use products in accordance with internal hospital guidelines.
1.4 Remove visible dirt with lint free nonwoven wipes wetted with sterile cold water (<40 °C, 104 °F)
1.5 Ensure that all surfaces are thoroughly wetted and keep moist for a minimum of 5 minutes, then repeat this step with another wipe.
1.6 Dispose protective gloves and disinfect your hands.

2. Intermediate level disinfection

2.1 Disinfect your hands and put on disposable gloves and proper protective gear.
2.2 Use disinfecting wipes* according the manufacturer’s instructions for use.
2.3 Let product work in for 5 minutes and wipe afterwards with another disinfecting wipe*.
2.4 Allow the surface to dry for a minimum of 5 minutes.
2.5 Dispose protective gloves and disinfect your hands.

*Recommended agent for intermediate level disinfection:
CaviWipes®
Metrex® Research
Address: 1717 West Collins Avenue
Orange, CA 92867, USA
Homepage: http://www.metrex.com
Phone: +1 800 841 1428
E-Mail: metrexcustcare@sybrondental.com
Mikrozid® AF Wipes,
Schülke & Mayr GmbH
Address: Robert-Koch-Str. 2,
22851 Norderstedt, DEUTSCHLAND
Homepage: http://www.schuelke.com
Telefon: +49 40 521 00 0
E-Mail: info@schuelke.com

3. Storage

3.1 After appropriate cleaning and disinfecting check the device before re-use.
3.2 Store products dry and dust free under storing conditions (see chapter 14  Technical specifications).
11 Warranty, maintenance and checks

Warranty
Medaxis AG warrants the device will be free from defects in materials and workmanship for a period of 2 years from the date of delivery ex works. Faulty material will be replaced free of charge during this period if not resulting from abuse or misapplication. This will not apply to parts subject to wear and tear in use. To ensure compliance with this warranty as well as optimum service from Medaxis products, we recommend the exclusive use of Medaxis accessories with our appliances. In no event shall Medaxis AG be liable for claims which exceed the scope of warranty described including liability for consequential damages, etc. The right to the replacement of faulty parts will not be recognized by Medaxis if any work has been carried out on debritom® by unauthorized persons. This warranty is subject to the appliance being returned to a Medaxis service centre.

Maintenance
Maintenance according to EN/IEC 62353
Medaxis recommends to performing an inspection of the debritom+ once a year. The inspection needs to be carried out according to EN/IEC 62353 and should include the following procedures:
- Visual examination of the device, in particular silicone sealing ring
- Measurement of grounding conductor according EN/IEC 62353, measure at the metal piece will be seen after the pump has been removed (Note – do not measure at the two plungers!)
- Measurement of the insulation resistance according to EN/IEC 62353, the pole for rinsing fluids is an example for an exposed conductive part (measurement should be performed at the threaded part of the knurled head screw).
- Measurement of leakage current according to EN/IEC 62353 is inapplicable. Alternatively, the earth leakage current could be measured according to IEC 60601-1:2005
- Function control: the power consumption should not exceed the value given on the type label – measured with running motor at highest intensity level setting. As you remove the pump during operation (Note: pump could be damaged), there has to be an error message displayed and the device will be shut down immediately.

For any test described above, there is no need to open the device.

Examine the complete device on the outside for any cracks in the housing, and replace the silicone sealing ring as you should discover wear and tear or brittleness

Replacement with new silicone sealing ring

**Only in case of broken sealing ring**

Installation or replacement of battery

If the foot on/off switch needs a battery installation or replacement, turn on debritom® by pressing the ON button, and follow the instructions below:

1. Remove protective cover from foot on/off switch as shown (phillips screw driver needed)

2. Open battery cover (screwdriver). Insert or replace batteries and lock the battery cover. Make sure the blue LED lights up as you activate the foot on/off switch. Battery type - Alkali AAA (LR03) 1.5V. Do not use lithium batteries or rechargeable batteries.

Reassemble protective cover in reverse order.
Pairing foot on/off switch with debritom®

The foot on/off switch needs to be paired only in case of replacement. Turn on debritom® by pressing the on/standby button, and follow the instructions below:

1. Remove protective cover from foot on/off switch as shown (phillips screw driver needed)

2. Simultaneously press the on/standby and „+” buttons for > 1 second on debritom®

3. As soon as the symbol ±± appears, press the yellow button on the foot pedal within 10 seconds.

4. As soon as the foot on/off switch has been detected by the device ±± press the yellow button within 5 seconds on the foot on/off switch again to complete the pairing.

5. If the pairing was successful, the following symbol ±± appears on the display - continue with step 6.
If the pairing was not successful, press the yellow button on the foot pedal for more than 7 seconds (existing pairing will be deleted) and repeat steps 2 to 5.

6. Reattach protective cover

Safety check

There are no prescribed or recommended safety checks to be carried out. Each time the device is switched on, a self-test is carried out which checks the internal functions of debritom®.

Due to the device layout and construction design, Medaxis does not expect electrical safety to be affected at any time during the life of the product - provided that debritom® is repaired throughout its service life strictly and exclusively by Medaxis authorized service centers and that it is used properly in accordance with the intended use.

The safety instructions must be observed.

12 Disposal

debritm® comprises metals and plastics and should be disposed of in accordance with the European directives 2002/95/EC and 2002/96/EC. Additional, local guidelines must also be observed. The electronic components must be disposed of separately, in accordance with the local regulations. Please take care that you dispose of debritom® and its accessories in accordance with the hospital disposal guidelines.

User information for the disposal of electrical and electronic equipment

This symbol means that the electrical and electronic equipment must not be disposed household refuse. A correct disposal of this device protects and prevents possible damage to the environment or human health. For more information about the disposal contact the manufacturer, your local caregiver or healthcare provider.

This symbol is only valid in the European Union. Please respect the relevant state laws and rules in your country for the disposal of electrical and electronic equipment.

13 Accessories

 WARNINGS

debritm® was validated in combination with the accessories listed in Chapter 17. For a correct and safe operation use debritom® with these accessories only. Further information is supplied with the individual accessory.
14 Technical Specifications

5.3kg/11.7lbs

[VAC] 100 – 240
[Hz] 50/60
[VA] 200

HxWxD
44x25x24 cm
18x10x10*

Transport / Storage
Humidity 15 – 93%
Temperature 5 – 40°C / 41 – 104°F

Operation
Humidity 15 – 93%
Temperature 5 – 37°C / 41 – 98°F

Fuse: Schurter AG, P/N: 0001.2512, SPT 5 x 20 mm, T 6.3AH, 250 VAC

Operation below 2000 m (to sea level)

Classification according to 60601-1
– Medical equipment class I, external power supply
– Continuous operation
– Protection class IP20
– The Handpiece is a applied part type B as a possible conductive water jet will touch the patient

15 Signs and symbols


This symbol indicates the legal specifications of the system.

This symbol indicates that the product or company has successfully met stringent standards for product safety.

This symbol indicates that interferences may occur in the vicinity of equipment marked with this symbol.

This symbol indicates the date of manufacture (four digits for the date).

This symbol indicates the manufacturer.

This symbol indicates the temperature limit –ation for operation, transport and storage.

This symbol indicates the humidity limitation for operation, transport and storage.

This symbol indicates the device is sterilized using ethylene oxide.

This symbol indicates the device is a single use device. Do not reuse the device.

This symbol indicates MR UNSAFE

This symbol indicates that the device should not be used after the end of the date shown.

This symbol indicates the device should not be used after the end of the date shown.

This symbol indicates the device should not use the device if package is damaged.

This symbol indicates the device should not use the device if package is damaged.

This symbol indicates the temperature limitation for operation, transport and storage.

This symbol indicates the number of pieces per package.

This symbol indicates a prescription device. Federal law restricts this device to sale by or on the order of a licensed healthcare practitioner (for US only).

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This symbol indicates a single use device. Do not reuse the device.

This symbol indicates that interferences may occur in the vicinity of equipment marked with this symbol.

This symbol indicates the date of manufacture (four digits for the date).

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This symbol indicates that the device should not be used after the end of the date shown.

This symbol indicates do not use the device if package is damaged.

This symbol indicates do not use the device if package is damaged.

This symbol indicates the number of pieces per package.

This symbol indicates the number of pieces per package.

debritom® handpiece standard jet

debritom® handpiece flat jet wide

Indicates the item is a medical device
Symbols for the error messages are described in chapter 9, 'Error messages'.

**Search for Bluetooth device for pairing**
Pressing the on/standby and ,+ buttons for > 1 s at debritom® will show the 'Search for Bluetooth device' symbol for pairing a new or replacement of the foot on/off switch.

**Bluetooth signal of foot on/off switch detected - confirm**
As soon as debritom® has detected the Bluetooth signal of the foot on/off switch, this symbol appears on the display.

**debritom® connected with foot on/off switch**
This symbol appears on the display as the foot on/off switch is connected to debritom®.

**Intensity levels**
The intensity levels can be adjusted between 1 - 5 (1 weakest/5 strongest)

--

16 Technical Documentation

**EMC**
debritom® is EMC-tested in conformity with the requirements of IEC 60601-1-2:2014 and can be used in the vicinity of other EMC-tested devices that fulfill the requirements of the relevant IEC 60601-1-2 standard. Untested HF (high-frequency) sources, radio networks or the like can influence the operation of the device and may not be operated in combination with the system. Debritom® is a medical device that requires special safety precautions and must be installed and placed in operation in accordance with the attached EMC information. Portable and mobile RF communication devices (mobile telephones) can affect debritom®.

The debritom® system is suitable for use in hospital environments, except for near active HF Surgical Equipment and except for RF shielded room of a magnetic resonance imaging system.

The debritom® does not provide any Essential Performance according to IEC 60601-1. The EMC immunity was tested in the "Ready to run" operating mode, the main focus of the immunity testing was to avoid an unintended start of water jet.

The following precautions should be taken to prevent adverse events to the Patient and Operator due to electromagnetic disturbances:

- Mains power quality should be that of a typical commercial and/or hospital environment.
- Floor should be wood, concrete, or ceramic tile. If floors are covered with synthetic material, the relative humidity should be at least 10%.
- Power frequency magnetic fields should be at levels characteristic of a typical location in a typical commercial or hospital environment.

**WARNING**
Use of this equipment adjacent to or stacked with other equipment should be avoided because it could result in improper operation. If such use is necessary, this equipment and the other equipment should be observed to verify that they are operating normally Portable and mobile RF communications equipment can affect medical electrical equipment.

Cables and accessories not specified within this user manual are not authorized. Using other cables and/or accessories may adversely impact safety, performance and electromagnetic compatibility (increased emission and decreased immunity).

**WARNING**
Use of accessories, transducers and cables other than those specified or provided by the manufacturer of this equipment could result in increased electromagnetic emissions or decreased electromagnetic immunity of this equipment and result in improper operation.

**WARNING**
Portable RF communications equipment (including peripherals such as antenna cables and external antennas) should be used no closer than 30 cm (12 inches) to any part of the Debritom® System, including cables specified by the manufacturer. Otherwise, degradation of the performance of this equipment could result.

To maintain basic safety and essential performance in regards to EMC no service is necessary.
The debritom+ System was tested according to IEC 60601-1-2:2014 with the following limits:

<table>
<thead>
<tr>
<th>Emissions Test</th>
<th>Compliance</th>
</tr>
</thead>
<tbody>
<tr>
<td>RF emissions</td>
<td>Group 1</td>
</tr>
<tr>
<td>CISPR 11</td>
<td>Class B</td>
</tr>
<tr>
<td>Harmonic emissions</td>
<td>Class A</td>
</tr>
<tr>
<td>IEC 61000-3-2</td>
<td></td>
</tr>
<tr>
<td>Voltage fluctuations /</td>
<td>Complies</td>
</tr>
<tr>
<td>flicker emissions</td>
<td></td>
</tr>
<tr>
<td>IEC 61000-3-3</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Immunity Test</th>
<th>Test Level</th>
</tr>
</thead>
<tbody>
<tr>
<td>Electrostatic discharge (ESD)</td>
<td>±5 kV contact</td>
</tr>
<tr>
<td>IEC 61000-4-2</td>
<td>±5 kV air</td>
</tr>
<tr>
<td>Electrical fast transient/burst</td>
<td>±2 kV</td>
</tr>
<tr>
<td>IEC 61000-4-6</td>
<td>100 kHz repetiton frequency</td>
</tr>
<tr>
<td>Surge</td>
<td>±1 kV line to line</td>
</tr>
<tr>
<td>IEC 61000-4-5</td>
<td>±2 kV line to earth</td>
</tr>
<tr>
<td>Voltage drop, short interruptions and voltage variations on power supply input lines</td>
<td>±0.1 % Un for 0.5 cycle</td>
</tr>
<tr>
<td>IEC 61000-4-11</td>
<td>70 % Un for 25 cycles</td>
</tr>
<tr>
<td>Power frequency (50/ 60 Hz)</td>
<td>50 A/m</td>
</tr>
<tr>
<td>Magnetic field IEC 61000-4-8</td>
<td></td>
</tr>
<tr>
<td>Radiated RF EM fields IEC 61000-4-3</td>
<td>3 V/m</td>
</tr>
<tr>
<td></td>
<td>80 MHz - 2 THz</td>
</tr>
<tr>
<td></td>
<td>80% AM at 1 kHz</td>
</tr>
<tr>
<td>Proximity fields from RF wireless communications equipment:</td>
<td></td>
</tr>
<tr>
<td>385 MHz 27 V/m</td>
<td>450 MHz 28 V/m</td>
</tr>
<tr>
<td>715 MHz 9 V/m</td>
<td>765 MHz 5 V/m</td>
</tr>
<tr>
<td>780 MHz 9 V/m</td>
<td>810 MHz 8 V/m</td>
</tr>
<tr>
<td>810 MHz 26 V/m</td>
<td>870 MHz 28 V/m</td>
</tr>
<tr>
<td>930 MHz 9 V/m</td>
<td>1720 MHz 28 V/m</td>
</tr>
<tr>
<td>1725 MHz 28 V/m</td>
<td>1865 MHz 28 V/m</td>
</tr>
<tr>
<td>1970 MHz 28 V/m</td>
<td>2450 MHz 28 V/m</td>
</tr>
<tr>
<td>2450 MHz 28 V/m</td>
<td>5200 MHz 9 V/m</td>
</tr>
<tr>
<td>5500 MHz 9 V/m</td>
<td>5785 MHz 9 V/m</td>
</tr>
</tbody>
</table>

RF Transmitter/Receiver used in the equipment:

Bluetooth 4.0 (single mode/Bluetooth smart): max RF output power 3.2 mW 2.402-2.480 GHz, ISM band. RFID 13.56 MHz, phase jitter modulation, max power: 200 mW

Service:
Requirements for service personnel: The service personnel must be trained by Medaxis AG in order to service the debritom+ and its accessories.
Medaxis AG will make available on request circuit diagrams, component part lists, descriptions, calibration instructions, or other information for Medaxis authorized service centers.

17 Required and optional accessories

**Required accessories**

**Pumps**

- debritom+ pump REF 2000.0200
- debritom+ pump, one day (contains pump and luer cap) REF 2000.0201

**Handpiece**

- debritom+ handpiece standard jet REF 2000.0003
- debritom+ handpiece flat jet narrow REF 2000.0004
- debritom+ handpiece flat jet wide REF 2000.0005

**Connecting tube**

- debritom+ connecting tube REF 2000.0301

**Optional accessories**

- AeroGuard 'Jellyfish' REF 2000.2000
- DebriClip REF 2000.4001
To install debritom⁺, follow the steps below

Tool for attaching: hex wrench 6mm

Connection debritom⁺/trolley
Rail

Storage for accessories
Antistatic castors with brakes

Spare parts

<table>
<thead>
<tr>
<th>Item</th>
<th>Description</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>debritom⁺ power cord (US) L= 5m</td>
<td>REF 2000 5008</td>
<td></td>
</tr>
<tr>
<td>Knurled head screw for pole rinsing fluid</td>
<td>REF 8000 0000</td>
<td></td>
</tr>
<tr>
<td>Pole for rinsing fluid</td>
<td>REF 8000 0001</td>
<td></td>
</tr>
<tr>
<td>Seal drive unit</td>
<td>REF 8000 0002</td>
<td></td>
</tr>
<tr>
<td>Transportation safety bolt</td>
<td>REF 8000 0005</td>
<td></td>
</tr>
<tr>
<td>Foot on/off switch</td>
<td>REF 2000 5020</td>
<td></td>
</tr>
</tbody>
</table>

Rinsing fluid

The debritom⁺ system was validated with the rinsing fluids listed below: NaCl 0.9% and ringer solution.

There have been other rinsing fluids tested – please do not hesitate to contact us for further information.

Other rinsing fluids, antimicrobial fluids, disinfecting agents can be used as per prescription by a medical professional.

Should any questions regarding the use of other fluids arise, please contact the manufacturer of the rinsing solution.

SAFETY INSTRUCTION

Medaxis’ purpose of testing a variety of rinsing fluids is to make sure that these fluids work properly with debritom⁺, not causing any harm to the system.

In order to make sure the fluids you intend to use do not cause any harm to the patient, you must contact the manufacturer of the rinsing fluid.

The manufacturer of the rinsing fluid must be contacted for questions regarding medical application.

The debritom⁺ System was tested with bottles and bags up to 1000 ml.