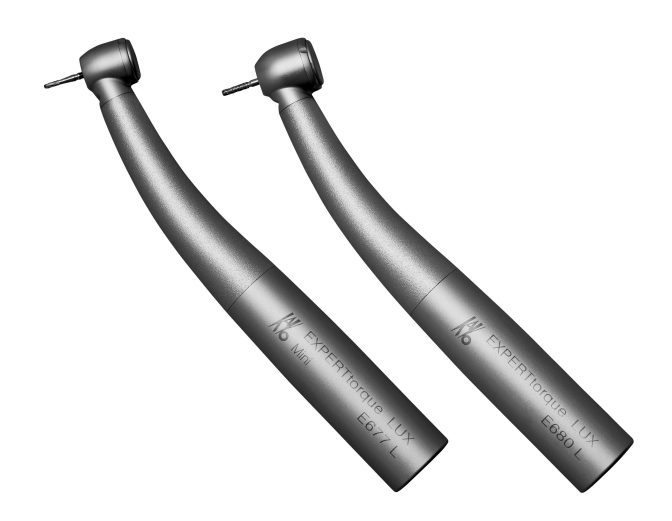
# Instructions for use

EXPERTtorque E680 - 1.006.8700, 1.006.4700, 1.006.4300, 1.006.9000, 1.006.4600 EXPERTtorque Mini E677 - 1.007.3600, 1.006.0100, 1.006.3800







KaVo Dental GmbH Bismarckring 39 D-88400 Biberach Phone +49 7351 56-0 Fax +49 7351 56-1488

## Manufacturer:

Kaltenbach & Voigt GmbH Bismarckring 39 D-88400 Biberach www.kavo.com



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1 User instructions

#### 1 User instructions

Dear User

Congratulations on purchasing this KaVo quality product. By following the instructions below you will be able to work smoothly, economically and safely.

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#### **KaVo Technical Service**

If you have any questions or complaints, please contact the KaVo Technical Service: +49 (0) 7351 56-1000 service.instrumente@kavokerr.com

## **KaVo Repair Service**

For repairs, please contact your local dealer or the KaVo Repair Service directly: +49 (0) 7351 56-1900 service.reparatur@kavokerr.com

## **Target group**

This document is intended for dentists and their assistants. The startup section is also intended for service technicians.

## **General marks and symbols**

| <u>^!\</u> | Refer to the chapter on Safety/Warning symbol  |
|------------|--|
| i          | Important information for users and service technicians  |
| <b>•</b>   | Action request   |
| CE         | CE mark (European Community). A product bearing this mark meets the requirements of the applicable EC directive. |
| 135°C      | Can be steam-sterilised at 134 °C -1 °C / +4 °C (273 °F -1.6 °F / +7.4 °F)                                       |
| X          | Thermodisinfectable  |

## Information on the packaging

| REF | Material number   |
|-----|---|
| SN  | Serial number   |
|     | Legal Manufacturer                                      |
|     | CE mark according to Medical Devices Directive EC 93/42 |

1 User instructions

| (i)  | Please note the electronic instructions for use           |
|------|---|
| Ŵ    | Note: observe accompanying documents                      |
| EAC  | EAC conformity mark (Eurasian Conformity)                 |
|      | GOST R certification                                      |
| oc C | Transportation and storage conditions (Temperature range) |
| hPa  | Transportation and storage conditions (Air pressure)      |
|      | Transportation and storage conditions (Humidity)          |
|      | Protect from moisture                                     |
| T    | Protect from impact                                       |
|      | HIBC Code   |

#### **Hazard levels**

The warning and safety notes in this document must be observed to prevent personal injury and material damage. The warning notes are designated as shown below:



## **⚠** DANGER

In cases which – if not prevented – directly lead to death or severe injury.



## **⚠** WARNING

In cases which – if not prevented – could lead to death or severe injury.



## **A** CAUTION

In cases which – if not prevented – could lead to minor or moderate injury.

## **NOTICE**

In cases which - if not prevented - could lead to material damage.

2 Safety | 2.1 Infection hazard

## 2 Safety

The instructions for use are a component of the product and must be read carefully prior to use and be accessible at all times.

The device may only be used in accordance with the intended use, any other type of use is not permitted.

#### 2.1 Infection hazard

Patients, users or third parties could be infected by contaminated medical devices.

- ► Take suitable personal protective measures.
- Follow the instructions for using the components.
- ▶ Before the initial startup and after each use, reprocess and sterilise the medical device and accessories accordingly.
- ► Carry out the cleaning and sterilisation as described in the instructions for use. The procedure has been validated by the manufacturer.
- ► It is essential to ensure the effectiveness of the cleaning and sterilisation in the case of deviation in procedure.
- Prior to disposal, the product and accessories must be appropriately reprocessed or sterilised.
- ► In the case of injury to soft tissue, do not continue treatment in the oral cavity with compressed air-driven instruments.

### 2.2 Technical condition

A damaged device or components could injure patients, users and third parties.

- ▶ Only operate devices or components if they are undamaged on the outside.
- ► Check that the device is working properly and is in satisfactory condition before each use.
- Have parts with sites of breakage or surface changes checked by the Service.
- ► If the following defects occur, stop working and have the service personnel carry out repair work:
- Malfunctions
- Damage
- Irregular running noise
- Excessive vibration
- Overheating
- Dental bur or diamond grinder is not firmly locked in the handpiece

Observe the following instructions in order to guarantee optimum functioning and prevent material damage:

- ► Service the medical device with care products and systems regularly as described in the instructions for use.
- ► The device should be cleaned, serviced and stored in a dry location, according to instructions, if it will not be used for a longer period.

2 Safety | 2.3 Accessories and combination with other equipment

## 2.3 Accessories and combination with other equipment

Use of un-authorised accessories or un-authorised modifications of the device could lead to injury.

- Only use accessories that have been approved for combination with the product by the manufacturer.
- ▶ Only use accessories that are equipped with standardised interfaces.
- ▶ Do not make any modifications to the device unless these have been approved by the manufacturer of the product.

## 2.4 Qualification of personnel

Application of the product by users without the appropriate medical training could injure the patients, the users or third parties.

- ► Make sure that the user has read and understood the instructions for use.
- ► Only employ the device if the user has the appropriate medical training.
- Observe national and regional regulations.

The improper use of the device could lead to burns or injuries.

- ▶ Never touch soft tissue with the handpiece head or instrument cover.
- ▶ Do not use the medical device as a light probe.
- ► Use an appropriate light probe for illumination of the oral cavity or site of preparation.
- ► After treatment, place the medical device properly in the holder, without the cutter or grinder.

## 2.5 Service and repair

Repairs, servicing and safety checks may only be performed by trained service personnel. The following persons are authorised to do this:

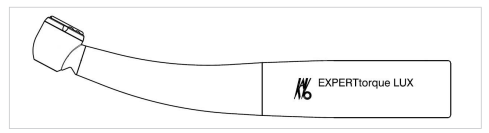
- Service technicians of KaVo branches after the appropriate product training
- Service technicians of KaVo authorised dealers after the appropriate product training

Observe all the following items during servicing work:

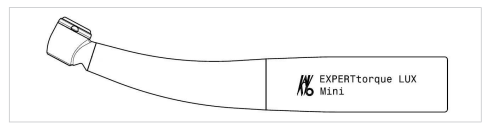
- ► Have the service and testing tasks carried out according to the Medical Product Operator Ordinance.
- ► After servicing, interventions, and repairs of the device and before re-use, the device must be subjected to safety checks by the service personnel.
- Following expiry of the warranty, have the tool holding system checked once a year.
- KaVo recommends specifying in-house service intervals where the medical device is brought to a professional shop for cleaning, servicing and a function check. Defined the service interval depending on the frequency of use.

3 Product Description | 3.1 Intended use

## **3 Product Description**



- EXPERTtorque LUX E680 L Mat. no. 1.006.8700
- EXPERTtorque E680 C Mat. no. 1.006.9000
- EXPERTtorque LUX E680 LN Mat. no. 1.006.4700
- EXPERTtorque LUX E680 LM Mat. no. 1.006.4300
- EXPERTtorque LUX E680 LS Mat. no. 1.006.4600



- EXPERTtorque Mini LUX E677 L Mat. no. 1.007.3600
- EXPERTtorque Mini LUX E677 LN Mat. no. 1.006.0100
- EXPERTtorque Mini LUX E677 LM Mat. no. 1.006.3800

#### 3.1 Intended use

#### Indications for use:

This medical device is

- intended for dental treatment only. All other types of use or modifications of the product are not permitted and can be hazardous. The medical device is intended for the following applications: Removal of carious material, cavity and crown preparations, removal of fillings, processing of tooth and restoration surfaces.
- A medical device according to relevant national statutory regulations.

#### **Proper use:**

According to these provisions, this medical device may only be used for the described application by a properly trained user. You need to comply with the following:

- the applicable health and safety regulations
- the applicable accident prevention regulations
- these Instructions for use

According to these regulations, the user is required to:

- to only use equipment that is operating correctly,
- adhere to the specified intended use
- to protect him or herself, the patient and third parties from hazards, and
- to prevent contamination from the product

3 Product Description | 3.2 Technical Specifications

## 3.2 Technical Specifications

|                                  | KaVo MUL-<br>TIflex |           | Morita<br>Alpha Con-<br>nection | Sirona R/F |
|----------------------------------|---------------------|-----------|---------------------------------|------------|
| Drive pressure (bar)             | 2.1 - 3.5           | 2.0 - 3.0 | 3.5 - 3.7                       | 2.6 - 3.0  |
| Drive pressure recommended (bar) | > 2.8               | 2.5       | 3.6                             | 2.7        |
| Return air pressure (bar)        | < 0.5               | < 0.5     | < 0.3                           | < 0.3      |
| Spray water pressure (bar)       | 0.8 - 2.0           | 0.8 - 2.0 | 0.5 - 2.0                       | 0.8 - 2.2  |
| Spray air pressure (bar)         | 1.0 - 2.5           | 1.0 - 2.5 | -                               | 0.8 - 2.9  |

|   | KaVo MUL-<br>TIflex                  | _ | Morita<br>Alpha Con-<br>nection | Sirona R/F |
|---|--------------------------------------|---|---------------------------------|------------|
| Air consumption (NI/min) 39 - 51                              |                                      |   |                                 |            |
| Idling speed (rpm) EX-<br>PERTtorque / EXPERT-<br>torque Mini | 340,000 - 420,000 / 400,000- 480,000 |   |                                 |            |
| Recommended application force (N)                             | 2 - 3                                |   |                                 |            |

## **Note**

KaVo MULTIflex is a registered trademark of Kaltenbach & Voigt GmbH, Biberach.

NSK PTL - CL- LED 3 is a product name of NSK NAKANISHI INC., Japan. Morita Alpha Connection is a product name of MORITA Dental Company, Japan.

Sirona is a registered brand of the company Sirona Dental System GmbH, Bensheim.

## 3.3 Transportation and storage conditions

#### **NOTICE**

## Startup after refrigerated storage.

Malfunction.

▶ Prior to startup, strongly refrigerated products must be allowed to warm up to a temperature of 20 °C to 25 °C (68 °F to 77 °F).

| -20°C    | Temperature: -20°C to +70°C (-4°F to +158°F)               |
|----------|--|
| 5%       | Relative humidity: 5% RH to 95% RH absence of condensation |
| 1060hPa  | Air pressure: 700 hPa to 1060 hPa (10 psi to 15 psi)       |
| <b>*</b> | Protect from moisture                                      |



4 Start up and shut down | 4.1 Checking the amount of water

## 4 Start up and shut down



## **MARNING**

#### Hazard from non-sterile products.

Infection hazard for dentist and patient.

 Prior to initial startup and after each use, reprocess the product and accessories.

#### See also:

₱ 7 Reprocessing steps in accordance with ISO 17664, Page 16



#### **MARNING**

## Dispose of the product in appropriate manner.

Infection hazard.

▶ Reprocess and sterilise the product and accessories before disposal.

#### See also:

₱ 7 Reprocessing steps in accordance with ISO 17664, Page 16

#### **NOTICE**

## Damage from soiled and moist cooling air.

Contaminated and moist cooling air can cause malfunctions.

▶ Make sure that the supply of cooling air is dry, clean, and uncontaminated according to EN ISO 7494-2.

## 4.1 Checking the amount of water

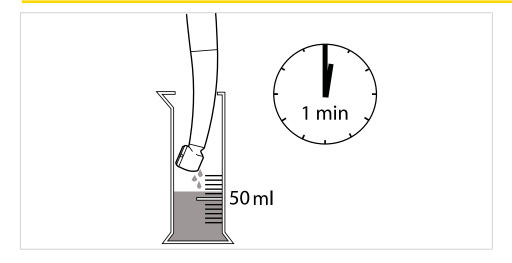






Overheating of the tooth due to insufficient amount of cooling water. Insufficient spray water can cause the medical device to overheat and damage the pulp and tooth.

- ► Adjust the water amount for the spray cooling to a minimum of 50 cm3/min (3.1 inch³).
- ► Check the spray water channels and clean the spray nozzles with the nozzle needle **Mat. no. 0.410.0921**.



4 Start up and shut down | 4.2 Checking the pressures

## 4.2 Checking the pressures

## **NOTICE**

# Contaminated or moist compressed air at the compressed air connection

Premature wear

► Ensure that the cooling air is dry, clean and uncontaminated in accordance with EN ISO 7494-2.



- ► Insert the test manometer (Mat. no. 0.411.8731) between the coupling and the medical device and check the following pressures:
- ⇒ Drive pressure, drive pressure recommended, return air pressure, spray water pressure, and spray air pressure.

#### See also:

3.2 Technical Specifications, Page 9

A higher drive pressure will be reduced automatically by the medical device.

5 Operation | 5.1 Attaching the medical device

## 5 Operation



#### **Note**

At the beginning of each workday, the water-conducting systems should be rinsed for at least 2 minutes (without transmission handpieces being attached) and if there is a risk of contamination from reflux or back suction, the system may also need to be rinsed for 20 to 30 seconds after each patient.

## 5.1 Attaching the medical device



## **MARNING**

## Detachment of the medical device during treatment.

A medical device that is not properly locked can detach from the coupling during treatment.

Before each use, check if the medical device is securely locked onto the coupling.

#### **NOTICE**

Inaccurate coupling can destroy the high-pressure lamp or the LED of the coupling or reduce its service life.

Make sure that the turbine is accurately coupled and firmly seated on the coupling.



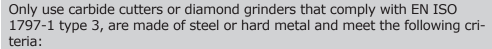
- Accurately attach the medical device to the coupling and push is to the rear until the coupling audibly locks in the medical device.
- ▶ Pull on it to make sure that the medical device is securely affixed to the coupling.

## 5.2 Removing the medical device

• Grasping the coupling, twist the medical device slightly and pull it off.

## 5.3 Inserting the milling cutter or diamond grinder

## Note





- Shaft diameter: 1.59 to 1.60 mm (0.0626 in to 0.0629 in)
- Overall length: EXPERTtorque: max. 25 mm
- Overall length of EXPERTtorque Mini: max. 19 mm
- Shaft clamping length: EXPERTtorque: min. 11 mm
- Shaft clamping length EXPERTtorque Mini: min. 9 mm
- Blade diameter: max. 2 mm (0.0787 in)

## **MARNING**



## Use of unauthorised dental burs or diamond grinders.

Injury to the patient or damage to the medical device.

- ► Comply with the instructions for use and use the dental bur or diamond grinder properly.
- Only use dental burs or diamond grinders that do not deviate from the specified data.

5 Operation | 5.4 Removing the milling tool or diamond grinder



## CAUTION

Use of dental burs or diamond grinders with worn or damaged shafts. Risk of injury, tool may fall out during treatment.

▶ Never use dental burs or diamond grinders with damaged or worn shafts.



## CAUTION

Danger of injury from cutters or grinders.

Infections or cuts.

Wear gloves or finger stalls.



## **CAUTION**

#### Hazard from defective chuck system.

The cutter or grinder could fall out and cause injury.

▶ Pull on the dental burr or rips abrasives to check if the clamping system is functioning properly and that the tool is firmly clamped. Wear gloves or a thimble to check, insert, or remove the bits to prevent injury and infection.

#### NOTICE

Tool shaft slips inside the chuck due to excessive speed of the tool or abrupt engagement of the tool.

Material damage to tool shaft and chuck system, reduction of the service life of tool and chuck system.

Do not operate the tool at a higher speed than recommended by the manufacturer.

#### **NOTICE**

Use of dental burs or diamond grinders with worn or damaged shafts. Material damage to the chuck system, tool is difficult or impossible to remove from the chuck system.

▶ Do not use dental burs or diamond grinders with damaged or worn shafts.



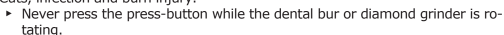
- Forcefully press the push button with your thumb and simultaneously insert the cutter or grinder all the way.
- Check that the cutter or grinder is seated by pulling on it.

## 5.4 Removing the milling tool or diamond grinder

#### ⚠ WARNING

Hazard from rotating dental bur or diamond grinder.

Cuts, infection and burn injury.



- Do not touch the dental bur or diamond grinder while it is rotating.
- Never touch soft tissue with the handpiece head or instrument cover.
- Remove the dental bur/diamond grinder from the contra-angle handpiece after treatment to avoid injury and infection when putting it away.

#### **NOTICE**

#### Damage to the chucking system.

Property damage.

► Do not press the pushbutton while the cutter or grinder is rotating.



5 Operation | 5.4 Removing the milling tool or diamond grinder



After the dental bur or diamond grinder has stopped rotating, firmly press the press-button with your thumb and simultaneously pull out the dental bur or diamond grinder.

6 Troubleshooting | 6.1 Cleaning the spray nozzle

## 6 Troubleshooting

## 6.1 Cleaning the spray nozzle



## **MARNING**

## Hazard from nonsterile products.

Infection danger to the care provider and patient.

▶ Reprocess and sterilise the medical device properly before the next use.



## **A** CAUTION

#### Hazard from insufficient spray water.

Overheating of the medical device and damage to the tooth.

- ► Check the spray water channels and clean the spray nozzles with the nozzle needle **Mat. no. 0.410.0921** if necessary.
- ► Check the water filter and exchange if necessary.



## 6.2 Changing the water filter



## **MARNING**

#### Hazard from nonsterile products.

Infection danger to the care provider and patient.

▶ Reprocess and sterilise the medical device properly before the next use.



## **A** CAUTION

#### Hazard from insufficient spray water.

Insufficient spray water can cause the medical device to overheat and damage the tooth.

- Check the filter and exchange if necessary.
- Check the spray water channels and clean the spray nozzles with the nozzle needle Mat. no. 0.410.0921 if necessary.



 Unscrew the sleeve ① in counterclockwise direction from the insert ② and pull it off.



- Unscrew the water filter ③ with the wrench (Mat. no. 1.002.0321) and take it out
- ► Insert the new filter (Mat. no. 1.002.0271) and screw it in with the wrench
- ▶ Place the sleeve ① on the insert ②, and screw it tight in clockwise direction.

7 Reprocessing steps in accordance with ISO 17664 | 7.1 Preparations at the site of use

## 7 Reprocessing steps in accordance with ISO 17664

## 7.1 Preparations at the site of use



## **MARNING**

## Hazard from contaminated products.

Contaminated products are associated with an infection hazard.

► Take suitable personal protective measures.



## **WARNING**

## Sharp tool in the medical device.

Injury hazard from sharp and/or pointed tool.

- ► Remove the tool.
- ► Reprocess the medical device as soon as possible after treatment.
- ► The medical device must be dry when transported to reprocessing.
- ► To minimise the risk of infection during reprocessing, always wear protective gloves.
- Remove the tool from the medical device.
- ► Remove all residual cement, composite or blood immediately.
- ▶ Do not place in solutions or similar substance.

## 7.2 Manual Reprocessing



## **MARNING**

## Sharp tool in the medical device.

Injury hazard from sharp and/or pointed tool.

Remove the tool.

#### NOTICE

## Never reprocess this medical device in an ultrasonic device.

Malfunction and material damage.

Clean manually or in a washer disinfector only.

## 7.2.1 Manual external cleaning

Required accessories:

- Tap water 30°C ± 5°C (86°F ± 10°F)
- Brush such as a medium hard toothbrush



Brush off under flowing tap water.

## 7.2.2 Manual internal cleaning

Validated internal cleaning (removal of residual protein) can be accomplished with KaVo CLEANspray.

► Cover the medical device with the KaVo Cleanpac bag, and place it on the corresponding care adapter.

7 Reprocessing steps in accordance with ISO 17664 | 7.2 Manual Reprocessing

- ► Hold the can vertically.
- ▶ Press the spray button three times for 2 seconds each time.
- ► Remove the medical device from the spray attachment and let the cleanser act for 1 minute.

#### See also:

- ► If a manual external and internal disinfection do not follow directly, dry the medical device with KaVo DRYspray.

#### See also:

₱ 7.2.5 Manual drying, Page 18

#### **Note**

KaVo CLEANspray and KaVo DRYspray for manual interior cleaning are only available in the following countries:

Belgium, Denmark, Germany, Finland, France, United Kingdom, Italy, Latvia, Lithuania, Luxembourg, Netherlands, Norway, Austria, Poland, Portugal, Sweden, Switzerland, Spain and the Czech Republic.

In other countries interior cleaning can only be carried out with washer disinfectors in accordance with EN ISO 15883-1.

## 7.2.3 Manual external disinfection

#### **MARNING**

#### Incomplete disinfection.

Infection hazard.

- Only use disinfection procedures that are verified to be bactericidal, fungicidal and virucidal.
- ► If the disinfectants used do not meet these requirements, the process must be concluded by disinfection of the unit(s) without packaging using a steam steriliser.

#### **NOTICE**

## Never disinfect the handpiece with chloride-containing products.

Malfunction and material damage.

Only disinfect in a washer disinfector or manually.

KaVo recommends the following products based on compatibility of the materials. The microbiological efficacy must be ensured by the disinfectant manufacturer and proven by an expert opinion.

#### Approved disinfectants:

- Mikrozid AF made by Schülke & Mayr (liquid or cloths)
- FD 322 made by Dürr
- CaviCide made by Metrex

#### Consumables required:

- Cloths for wiping the medical device.
- Spray the disinfectant on a cloth, then wipe down the medical device and allow the disinfectant to act according to the instructions of the disinfectant manufacturer.







7 Reprocessing steps in accordance with ISO 17664 | 7.2 Manual Reprocessing

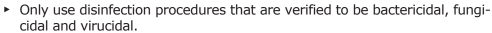
► Follow the instructions for use of the disinfectant.

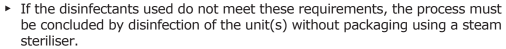
#### 7.2.4 Manual internal disinfection

## **MARNING**

## Incomplete disinfection.

Infection hazard.





#### NOTICE

**Never disinfect the handpiece with chloride-containing products.**Malfunction and material damage.

Only disinfect in a washer disinfector or manually.

The efficacy of manual internal disinfection must be demonstrated by the manufacturer of the disinfection agent. With KaVo products, use only disinfection agents that have been released by KaVo with respect to the compatibility of materials (e.g. WL-cid / made by ALPRO).

- ► Cover the medical device with the KaVo CLEANpac bag, and place it on the corresponding care adapter.
- Hold the can vertically.
- Press the spray key for at least 3 seconds.
- ► Remove the medical device from the spray attachment and let the disinfectant act for 2 minutes.
- ► Follow the instructions for use of the disinfectant.

## 7.2.5 Manual drying

Use KaVo DRYspray for subsequent drying of the air, water and gear unit ducts.

- ► Cover the medical device with the KaVo CLEANpac bag, and place it on the corresponding care adapter.
- Hold the can vertically.
- Press the spray key for at least 3 seconds.

#### See also:

KaVo DRYspray Instructions for Use

#### **Note**

KaVo CLEANspray and KaVo DRYspray for manual interior cleaning are only available in the following countries:

Belgium, Denmark, Germany, Finland, France, United Kingdom, Italy, Latvia, Lithuania, Luxembourg, Netherlands, Norway, Austria, Poland, Portugal, Sweden, Switzerland, Spain and the Czech Republic.

In other countries interior cleaning can only be carried out with washer disinfectors in accordance with EN ISO 15883-1.

► Immediately after drying, lubricate the KaVo medical device with care agents from the KaVo care system.



7 Reprocessing steps in accordance with ISO 17664 | 7.3 Automated Reprocessing

#### See also:

₱ 7.4 Care products and systems - Servicing, Page 20

## 7.3 Automated Reprocessing

## **MARNING**

#### Incomplete disinfection.

Infection hazard.

- Only use disinfection procedures that are verified to be bactericidal, fungicidal and virucidal.
- ▶ If the disinfectants used do not meet these requirements, the process must be concluded by disinfection of the unit(s) without packaging using a steam steriliser.



## **MARNING**

## Sharp tool in the medical device.

Injury hazard from sharp and/or pointed tool.

Remove the tool.

#### NOTICE

**Never disinfect the handpiece with chloride-containing products.**Malfunction and material damage.

► Only disinfect in a washer disinfector or manually.

#### **NOTICE**

Never reprocess this medical device in an ultrasonic device.

Malfunction and material damage.

► Clean manually or in a washer disinfector only.

# 7.3.1 Automated internal and external cleaning and internal and external disinfection



KaVo recommends washer disinfectors according to EN ISO 15883-1, which are operated using alkaline cleaning agents having a maximum pH value of 10. The validation was conducted with a Miele washer disinfector using the "VARIO-TD" programme, the "neodisher® mediClean" cleaning agent, the neodisher® Z" neutralisation agent and the "neodisher® mielclean" rinsing agent.

► For program settings as well as cleansers and disinfectants to be used, please refer to the Instructions for Use of the thermodisinfector.

## 7.3.2 Automated Drying

The drying procedure is normally part of the cleaning program of the thermodisinfector.



#### **Note**

Please observe the instructions for use of the thermodisinfector.

- ► In order to prevent impairment of the KaVo medical device, make sure that the inside and outside of the device is dry after the end of the cycle.
- Remove any residual liquids with KaVo DRYspray.

7 Reprocessing steps in accordance with ISO 17664 | 7.4 Care products and systems - Servicing

#### See also:

- ₱ 7.2.5 Manual drying, Page 18
- ► Immediately after drying, lubricate the KaVo medical device with care agents from the KaVo care system.

## 7.4 Care products and systems - Servicing



#### **MARNING**

#### Sharp tool in the medical device.

Injury hazard from sharp and/or pointed tool.

Remove the tool.



## **⚠** CAUTION

#### Improper service and care.

Risk of injury.

Service regularly with suitable agents.



#### Note

KaVo guarantees the proper function of KaVo products only if the care products listed as accessories are used, since these were tested for proper use on our products.

## 7.4.1 Servicing with KaVo Spray

KaVo recommends servicing the product as part of the reprocessing after each use, i.e. after each cleaning, disinfection, and before each sterilisation.

Remove the tool from the medical device.



- Cover the medical device with the KaVo CLEANpac bag, and place it on the corresponding care adapter.
- Press the spray key for 1 to 2 seconds.

#### **Chuck care**

KaVo recommends servicing the chucking system once weekly.

▶ Remove the tool from the medical device.



- Position the tip of the spray nipple in the opening, and apply the spray.
- ▶ Press the spray key for 1 to 2 seconds.

## 7.4.2 Servicing with KaVo QUATTROcare PLUS

Cleaning and servicing device with expansion pressure for internal cleaning of inorganic residues and optimum care.

(no validated cleaning of the interior according to German Robert Koch Institute (RKI) requirements)

KaVo recommends servicing the product as part of the reprocessing after each use, i.e. after each cleaning, disinfection, and before each sterilisation.

Remove the tool from the medical device.

7 Reprocessing steps in accordance with ISO 17664 | 7.4 Care products and systems - Servicing



Service the product in the QUATTROcare PLUS.

#### See also:

Instructions for use KaVo QUATTROcare PLUS

## Servicing of the clamping chuck

KaVo recommends servicing the chuck system once a week using the chuck servicing program integrated in the device.



#### **Note**

Handpieces must be taken off the service couplings before the chuck service can be started and performed.

- ► Close the front door and press the chuck service button for at least three seconds until the spray canister control LED flashes three times consecutively.
- □ The device is in chuck service mode.
- Remove the service coupling of the chuck from the side hatch of the QUAT-TROcare PLUS and attach it to coupling service point four, on the far right. A MULTIflex adaptor must be mounted there.



- Press the handpiece together with the guide bush of the chuck to be serviced against the tip of the service coupling.
- Press the button marked with the chuck service symbol.



#### Note

#### Close the chuck service mode.

Option 1: Place the dental handpieces in the QUATTROcare PLUS 2124 A, close the front door and start theservice procedure.

Option 2: After three minutes with no service procedure running, the device automatically switches back to normal service mode.

#### See also:

Servicing with KaVo QUATTROcare PLUS

## 7.4.3 Care with KaVo SPRAYrotor



#### **Note**

KaVo SPRAYrotor is no longer included in the current delivery programme.

Follow-up product:

► QUATTROcare PLUS 2124 A

KaVo recommends servicing the product as part of the reprocessing after each use, i.e. after each cleaning, disinfection, and before each sterilisation.



- Cover the medical device with the Cleanpac bag, and place it on the corresponding servicing adapter on the KaVo SPRAYrotor.
- Service the product.

7 Reprocessing steps in accordance with ISO 17664 | 7.5 Packaging

#### See also:

Instructions for use KaVo SPRAYrotor

## 7.4.4 Servicing with KaVo QUATTROcare

# $f{i}$

## **Note**

QUATTROcare 2104 / 2104 A is no longer included in the current delivery programme.

Follow-up product:

QUATTROcare PLUS 2124 A

Servicing and cleaning device with expansion pressure for the interior cleaning of inorganic residues and optimum care.

(no validated cleaning of the interior according to German Robert Koch Institute (RKI) requirements)

KaVo recommends servicing the product as part of the reprocessing after each use, i.e. after each cleaning, disinfection, and before each sterilisation.

Remove the tool from the medical device.



► Service the product in the QUATTROcare.

#### See also:

☐ Instructions for use KaVo QUATTROcare 2104 / 2104A / 2124A

## Servicing the clamping chuck

KaVo recommends servicing the chucking system once weekly.

▶ Remove the tool from the medical device.



- ► Plug the spray nipple of the chuck servicing set onto the QUATTROcare plus Spray.
- ► Position the tip of the spray nipple in the opening, and apply the spray.
- Press the spray key for 1 to 2 seconds.

## 7.5 Packaging



#### **Note**

The sterilisation bag must be large enough for the instrument so that the bag is not stretched.

The quality and use of the sterilisation packaging must comply with applicable standards and be suitable for the sterilisation procedure!

► The medical device must be packed before sterilisation.

7 Reprocessing steps in accordance with ISO 17664 | 7.6 Sterilisation

#### 7.6 Sterilisation

# Sterilisation in a steam steriliser (autoclave) in accordance with EN 13060 / ISO 17665-1



## **⚠** CAUTION

#### Improper service and care.

Risk of injury.

Service regularly with suitable agents.

#### **NOTICE**

#### Contact corrosion due to moisture.

Damage to product.

► Immediately remove the product from the steam steriliser after the sterilisation cycle.



The KaVo medical device has a maximum temperature resistance up to 138  $^{\circ}$ C (280.4  $^{\circ}$ F).

## **Sterilisation parameters:**

Select a suitable procedure (depending on the available autoclave) from the following sterilisation processes:

- Steriliser with triple pre-vacuum:
  - at least 3 minutes at 134 °C -1 °C / +4 °C (273 °F -1.6 °F / +7.4 °F)
- Steriliser using the gravity method:
  - at least 10 minutes at 134 °C -1 °C / +4 °C (273 °F -1.6 °F / +7.4 °F) or alternatively
- at least 60 minutes at 121 °C -1 °C / +4 °C (250 °F -1.6 °F / +7.4 °F)
- ► Remove contra-angle handpieces and turbine handpieces from the steam steriliser immediately after completion of the sterilisation cycle.
- Use according to the manufacturer's Instructions for Use.

# 7.7 Storage

Reprocessed products must be stored appropriately such that they are protected from germs (as far as possible) and dust, in a dry, dark, cool room.



#### **Note**

Comply with the expiry date of the sterilised items.

8 Optional aids

# 8 Optional aids

Available from dental suppliers.

| Material summary                                     | Mat. no.   |
|--|------------|
| Replacement turbine EXPERTtorque                     | 1.007.9313 |
| Replacement turbine EXPERTtorque Mini                | 1.007.9457 |
| Wrench for lid of EXPERTtorque                       | 0.411.3053 |
| Wrench for lid of EXPERTtorque Mini (new)            | 1.008.6133 |
| Wrench for lid of EXPERTtorque Mini (old)            | 1.006.3384 |
| Replacement filter                                   | 1.002.0271 |
| Wrench   | 1.002.0321 |
| Instrument stand 2151                                | 0.411.9501 |
| Insert for turbines                                  | 0.411.9902 |
| Nozzle pin   | 0.410.0921 |
| Cleanpac 10 units                                    | 0.411.9691 |
| KaVo MULTIflex spray head for KaVo Spray             | 0.411.9921 |
| NSK spray head for KaVo Spray                        | 1.005.8436 |
| Morita spray head for KaVo Spray                     | 1.005.8275 |
| SIRONA spray head<br>for KaVo Spray                  | 1.005.8365 |
| KaVo MULTIflex service coupling for KaVo QUATTROcare | 0.411.7991 |
| NSK service coupling for KaVo QUATTROcare            | 1.000.8786 |
| Morita service coupling for KaVo QUATTROcare         | 1.000.6063 |
| Sirona service coupling for KaVo QUATTROcare         | 1.000.7156 |

| Material summary                                     | Mat. no.   |
|--|------------|
| Adapter KaVo MULTIflex for KaVo CLEANspray/ DRYspray | 1.007.1775 |
| CLEANspray/ DRYspray Starter set 2116 P              | 1.007.0573 |
| KaVo CLEANspray 2110 P                               | 1.007.0579 |
| KaVo DRYspray 2117 P                                 | 1.007.0580 |
| KaVo Spray 2112 A                                    | 0.411.9640 |
| ROTAspray 2142 A                                     | 0.411.7520 |
| QUATTROcare plus Spray 2140 P                        | 1.005.4525 |
| Chuck servicing set                                  | 1.003.1253 |

9 Terms and conditions of warranty | 7.7 Storage

## 9 Terms and conditions of warranty

The following warranty conditions apply to this KaVo medical device:

KaVo provides the end customer with a warranty of proper function and guarantees zero defects in respect of material and workmanship for a period of 18 months from the date of the invoice, subject to the following conditions: In case of justified complaints, KaVo will honour its warranty with a free replacement or repair. Other claims of any kind whatsoever, in particular with respect to compensation, are excluded. In the event of default, gross negligence or intent, this shall only apply in the absence of mandatory legal regulations to the contrary.

KaVo shall not be liable for defects and their consequences that have arisen or may arise from natural wear, improper handling, cleaning or maintenance, non-compliance with operating, maintenance or connection instructions, calcination or corrosion, contaminated air or water supplies or chemical or electrical factors deemed abnormal or impermissible in accordance with KaVo's instructions for use or other manufacturer's instructions. The warranty granted does not usually extend to lamps, optical fibres made of glass and glass fibres, glassware, rubber parts, and the colourfastness of plastic parts.

All liability is excluded if defects or their consequences originate from manipulations or changes to the product made by the customer or a third party that is not authorised by KaVo.

Warranty claims will only be accepted if the product is submitted along with proof of purchase in the form of a copy of the invoice or note of delivery. The dealer, purchase date, type, and serial number must be clearly evident from this document.



