HELIODENT DS

Operating Instructions

English
General information

Dear customer,

Thank you for purchasing your HELIODENT DS.

Both intraoral and mandibular joint radiography can be performed with this unit.

The technical documentation included in delivery, i.e. operating instructions, installation instructions, Installation report / Warranty passport, customer installation description and X-ray system file (only in the FRG), are also an integral part of this product.

Keep these documents nearby for quick and easy reference at all times (in the FR of Germany: in your X-ray system file).

In order to protect your warranty rights, please complete the enclosed Installation report / Warranty passport, together with the service technician, immediately after installation of your system.

Read the Operating Instructions to familiarize yourself with the product before taking X-rays on the patient. Always be sure to observe the currently valid Radiation Protection Guidelines and the Safety Notes in these Operating Instructions.

The ordinance governing the use of radiation emitting equipment in the Federal Republic of Germany requires the operator to perform regular constancy tests to ensure the safety of operating personnel and patients.

Your HELIODENT team

Maintenance

To ensure the safety and health of patients, operators and third parties, inspections and maintenance calls must be carried out at the specified intervals in order to guarantee the operational safety and functional reliability of your product (IEC 601-1 / DIN EN 60601-1 etc.).

It is the responsibility of the operator to ensure that the required inspections and maintenance work are carried out.

In the event that the operator fails to fulfill the obligation to carry out inspections and maintenance or ignores error messages, Sirona Dental Systems GmbH and/or its authorized dealers can not assume liability for any damage attributable to this.

This product bears the CE label in accordance with the provisions of the Council Directive 93/42/EWG of June 14, 1993 concerning medical devices.
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1 Warning and safety information

Marking of warning and safety information

To prevent personal injury and damage to property, please observe the warning and safety information provided in these operating instructions. They are specially marked with a warning symbol and ATTENTION.

Symbols used

⚠️ Observe accompanying documents (on name plate)

Proper use

This unit is designed for the preparation of intraoral X-rays and X-rays of the mandibular joint. The unit must not be used in explosion hazard areas.

General safety information

Patient X-ray exposures may be performed only when the unit is functioning flawlessly. The patient must not be left unattended at the unit.

Modifications to the system

Modifications to this system which could impair the safety of operators, patients or third persons are prohibited by law!

For reasons of product safety, this product may be operated only with original Sirona accessories or third-party accessories expressly approved by Sirona. The user is responsible for dangers resulting from the use of non-approved accessories.

Disturbance of electronic equipment worn by the patient.

To prevent the malfunction of electronic devices and loss of stored data in connection with radio controlled clocks, telephone cards, etc. all such items must be removed from the patient prior to the X-ray exposure.

Radiation protection

The applicable radiation protection regulations must be observed. The prescribed radiation protection accessories must be used.

If the release button is equipped with a cable, the operator must leave the vicinity of the X-ray tube unit before triggering an exposure.

No one except the patient may remain in the exposure room. In exceptional cases a third party who is not a member of the practice staff may assist the patient. Visual contact with the patient and the unit must be maintained during every X-ray exposure.

In case of malfunction, interrupt the exposure immediately by letting go of the release button.
Electromagnetic compatibility (EMC)

The HELIODENT DS complies with the requirements of IEC 60601-1-2:2001.

Medical electrical devices are subject to special precautionary measures regarding EMC. They must be installed and operated as specified in the document "Installation Requirements".

Portable and mobile HF communication devices can influence medical electrical equipment. The use of mobile telephones in the practice or hospital area therefore must be prohibited.

Disposal

It applies generally that the national regulations have to be complied with when disposing of this product. Please observe the regulations applying in your country.

Within the European Economic Community the directive 2002/96/EEC (WEEE) for electrical and electronic devices requires environmentally compatible recycling / disposal.

Your product is marked with the adjacent symbol. With the goal of environmentally compatible recycling / disposal, your product must not be disposed of with the domestic refuse.

The black bar under the "refuse bin" symbol means that it has been put onto the market after 13.08.2005. (See EN 50419:2005)

Please note that this product is subject to the directive 2002/96/EEC (WEEE) and laws applicable in your country and must be sent for environmentally compatible recycling / disposal.

The X-ray tube assembly of this product contains a tube with a potential implosion hazard, a lead lining and mineral oil.

Please contact your dealer if your product should be finally disposed of.

Dismantling and reassembly

When dismantling and reassembling the unit, always proceed as outlined in the instructions for reassembly. This is necessary to ensure the unit’s proper functioning and stability.
2 Technical Description

Technical Data

Nominal voltage: 200 – 230V or 127V
Permissible line voltage fluctuations: + 6% – 10% (200 – 230V), ± 10% (127V)
Nominal current:

- at 230V: 4.6A
- at 127V: 11A

Rated line frequency: 50Hz / 60Hz
Line impedance:

- at 230V: 0.8 Ohm
- at 127V: 0.3 Ohm

House wiring fuse: 16A
Power requirement: < 1.5kW
Tube voltage: 60kV
Tube current: 7mA
High-voltage curve: Multipulse X-ray tube
Radiation times: 0.01 – 3.5 s
Mark-to-space ratio: 1:30 (radiation-time-dependent overheating protection)
Total filtration in X-ray tube unit:

- > 2mm Al / 70 kV IEC 60 522

Focal spot size acc. to IEC 336: Petrick P470/6.30/12G
Focal spot mark: SSD 8" (203mm) or 12" (305mm) = source-skin distance ≤ 60 mm

Diameter of radiation field at tube end:
Type of protection against electric shock:
Degree of protection against electric shock:

- Protection class I
- Type B

Degree of protection against penetration of water:

- Conventional unit (w/o protection against penetration of water)

Year of manufacture: (on the rating plate) 20xx
Operating mode: Continuous operation
Continuous power rating: 30W max
Rating (60kV, 7mA): 420W max
Anode material: Tungsten
Exposure data to determine leakage radiation:

- 0.25mA/60kV
- ≤ 0.25 mGy/h

Filament
Nominal line voltage: 6.6V
Nominal current: 1.6A


**Technical Data**

Nominal frequency: DC

Duty cycle: 1:30

Operating conditions (compliant with IEC 601-1)
- Ambient temperature: 10°C – 40°C (50°F – 104°F)
- Relative humidity: 30% – 75%

Transport and storage conditions (compliant with IEC 601-1)
- Temperature: -40°C – +70°C (-40°F – 158°F)
- Relative humidity: 10% – 95%

HELIODENT DS has been inspected by the VDE Testing and Certification Institute for compliance with:
- IEC 60 601-1
- IEC 60 601-2-28 / 1993
- IEC 60 601-1-3 / 1994
- IEC 60 601-2-7 / 1998

HELIODENT DS has been found to comply with these regulations.

The HELIODENT DS X-ray unit complies with AS / NZS 3200.1.0

Reg. No.: China

Reg. No.: China
SFDA (I) 20053301170

Original language of Operating Instructions: German
Illumination

Central ray

Anode angle 24° ± 1°

Focal spot marc

Sizes and Weight

No true scale.
Approx. length values.
Weight: approx. 3.4kg

132 102

161

64.5

169.3
Cool-down curve for X-ray tube housing

Cool-down curve for X-ray tube

Warm-up curve for X-ray tube housing
3 Controls and display elements

1. Power switch
2. Standby (ready) indicator lamp (LED)
3. X-radiation indicator lamp (LED)
4. Digital exposure time display
5. Object selector switch
6. Digital mode indicator lamp
7. Selector button: digital/conventional exposure
8. Release button
9. Scale for setting angle of inclination
10. X-ray field limitation
11. X-ray tube unit

- Mobile stand / Overhead model / Centro column / Split Concept: see page 20/21
4 Accessories

1. Test specimen for constancy test for conventional radiographic technique
   Order no. 59 69 779

2. Test specimen for constancy test for universal and full-size sensors
   Order no. 51 68 062

3. Tube extension to 12" SSD (30 cm)
   Order no. 33 14 742

4. Square cone extension on 12" SSD (30 cm)
   Order No. 58 61 468

5. X-ray field limitation blue with twist grip
   for Full Size sensor and for conventional radiographic technique 3x4 cm
   Order no. 46 81 974

6. X-ray field limitation black with twist grip
   for Universal sensor and for conventional radiographic technique 2x3 cm
   Order no. 60 00 579

7. Sensor holder set with sighting rings and guide rods
   manufactured by DENTSPLY RINN
   Sirona order no: 58 58 522

**NOTE**
Not all of the accessories shown here are included in the scope of delivery.
5 Preparing an exposure

5.1 Switching the unit ON

Set the main power switch to position I.

When doing so, be sure not to press the release button (8) or button D (7).

Following switch-on, a self-test executes. A short acoustic signal is then emitted and a 3-digit software status display appears. Then the standby indicator lamp (2) starts flashing and the selected exposure time lights up. After approx. 20 sec. have elapsed, the standby indicator lamp lights up continuously.

When the Digital mode indicator also lights up, the digital radiographic technique is selected (see next page).

The unit is ready for operation.

NOTE

If an error is detected during the self-test, a number ranging from 1 to 4 will appear on the digital display. (see chapter entitled “List of error codes”).

The LED (2) then flashes.

The unit is not ready for operation!

Switch the unit OFF and ON again.

NOTE

If the same error occurs again, please contact your service technician.
5.2 Selecting the radiographic technique

Digital

If you are working with a digital radiographic system (e.g. SIDEXIS), the Digital mode indicator lamp (6) on the unit must be lit up.

You can switch back and forth between the conventional and the digital radiographic technique by briefly holding down the D button. The resulting reduction in the exposure time will then be displayed.

Use an X-ray field limitation for the digital radiographic technique.

**NOTE**

To set the recommended exposure times for adults, always set the patient symbol to the highest exposure level within the relevant tooth symbol.

If you are working with children, always set to the lowest exposure level within the relevant tooth symbol!

Conventional

If you want to take conventional (film) exposures, the Digital mode indicator (6) must not be lit up.

You can switch back and forth between the conventional and digital radiographic technique by briefly pressing the D button. The Digital mode indicator (6) then switches off.

The exposure time for conventional exposures is then displayed.

Insert the X-ray field limitation for the conventional radiographic technique.
5.3 Basic setting at factory

**Digital with 8” tube**

- **0.05 s – 0.08 s**
  - 0.05 s
  - 0.06 s
  - 0.08 s

**Conventional with 8” tube**

- **0.32 s – 0.50 s**
  - 0.32 s
  - 0.40 s
  - 0.50 s

Example:

Exposure times with Class E film.
5.4 Adjusting the basic setting

The exposure times are factory set for the use of Class E film:

- e.g. Kodak Ekta Speed, Agfa-Dentus M2.

**NOTE**

*Digital and conventional exposure times are programmed separately. The factory adjusted DIGITAL programming is adjusted specifically for SIDEXIS.*

In case of varying exposure conditions (see below) the basic setting must be altered as follows:

**Example:**

Change over of the basic setting from Class E to Class D film for the conventional radiographic technique.

1. Set the large patient symbol to the center area of the object selector switch (5) (maxillary bicuspid/mandibular molar).

2. Press button D (7) (approximately 4 sec.) until the standby LED (2) and the Digital mode indicator (6) start flashing. Let go of button D.

3. By turning the object selector switch (5), you can now increase +, or decrease – the basic setting step-by-step (max. 8 steps).

4. Press the D button (7) to save the selected setting. The Ready LED (2) is lit up continuously.

5. Reset the object selector switch (5) to its original position. The basic setting is now displayed 3 steps higher.

Do not press the D button (7) if you do not want to save the setting. In this case, switch the unit OFF and then ON again.

**Varying exposure conditions**

- **E** for Class E films such as Kodak Ekta Speed and Agfa-Dentus M2
- **D** for Class D films such as Kodak Ultra Speed

**For Class D films:**

Set object selector switch (5) three steps higher.

**Film and developing conditions**

Due to varying film and developing conditions, it may be necessary to increase or decrease the exposure setting by one step.

Please refer to the Exposure times tables.
5 Setting the exposure data

Legend

Set the patient symbol for a
adult patient
or
child
to the tooth to be X-rayed.

Mandibular front tooth
bicusp / canine tooth
also used for bite-wing exposures

Maxillary front tooth

Maxillary bicuspid / canine tooth and mandibular molar

Maxillary molar

Occlusal exposure

3 exposure levels

For conventional X-rays only:

Each tooth symbol is divided into 3 exposure levels (exposure times) to suit stronger or weaker jaw regions.
The corresponding exposure time is digitally displayed.

Exposure times: see table.

Upper illustration:
8" SSD tube for conventional exposures
Class E film:
Adult, maxillary molar,
exposure time: 0.40 s.

Lower illustration:
8" SSD tube for conventional exposures
Class E film:
Adult, weak maxillary molar,
exposure time: 0.32 s.
Positioning the patient/x-ray tube unit

Adjust the X-ray tube unit

Have the patient sit down on the chair.

Right-angle technique (with X-ray field limitation)

The film or X-ray sensor is positioned with a holding system.

Sirona X-ray sensors may be used only with the holding systems recommended by us.

Always follow the Operating Instructions for intraoral radiography supplied with SIDEXIS.

45° angle technique (without X-ray field limitation)

The film or X-ray sensor must be held by the patient.

Angle of inclination

Exposure tube for the occlusal plane

Supramaxillary teeth

<table>
<thead>
<tr>
<th>Teeth Type</th>
<th>Angle</th>
</tr>
</thead>
<tbody>
<tr>
<td>Molars</td>
<td>35°</td>
</tr>
<tr>
<td>Bicuspsids and canine teeth</td>
<td>45°</td>
</tr>
<tr>
<td>Front teeth</td>
<td>55°</td>
</tr>
<tr>
<td>Occlusal exposure</td>
<td>80°</td>
</tr>
<tr>
<td>Occlusal exposure (inverted)</td>
<td>-90°</td>
</tr>
<tr>
<td>Front teeth</td>
<td>-20°</td>
</tr>
<tr>
<td>Bicuspsids and canine teeth</td>
<td>-10°</td>
</tr>
<tr>
<td>Molars</td>
<td>-5°</td>
</tr>
</tbody>
</table>

Submaxillary teeth

Bite-wing exposures are performed with the settings for submaxillary exposures.

**Example with 8" SSD tube for a conventional (film) exposure (Class E film):**

Adult,
tooth in bicuspid region,
exposure time: 0.20 s.

---

**Exposure of the mandibular joint**

Possible only in conventional exposure technique.

We recommend using 9x12cm or 13x18cm cassettes with intensifying screens (available from specialized dealers) for exposures of the mandibular joint.

The correct exposure time will vary according to the film-screen system used.
Releasing an exposure

CAUTION
Observe radiation protection regulations, see page 4.  
Check for correct exposure data.

- An exposure can be triggered by holding down the release button (8).

The X-RAY LED (3) remains lit up during the entire exposure.  
The exposure time is also indicated by an acoustic signal.  

During operation via remote control, the X-ray LED on the remote control is also lit up.  

- The exposure is finished when the radiation automatically switches off.

The Ready LED (2) flashes until the automatic tube cooling delay has expired (automatic exposure interlock).

Interrupting an exposure

The exposure can be interrupted by simply letting go of the release button.

The previously elapsed exposure time will then flash.  
The unit is again ready for operation following the automatic tube cooling delay.

If necessary repeat the exposure. For film exposures, first load a new film. For digital exposures, the system must first be in the Ready state.  

The unit may remain switched on until you have finished working.

Error message

If an error or fault is detected during an exposure, the exposure will be automatically interrupted. The error code will then appear on the digital display and the Ready LED (2) will simultaneously switch off.  

If error code 1 or 2 appears, please contact your local service technician.
10 Mobile stand / overhead model / Centro Column / Split Concept (optional)

**Mobile stand**

Use the two handles at the rear to move the mobile stand.

The mobile stand may be moved only **slowly** and over an **even** floor surface.

⚠️ **CAUTION**

Danger of injury.
Move the mobile stand only when the X-ray tube unit is in its home position and secured by the fastening strap (with support arm retracted ←).

The mobile stand has 4 casters with brakes.
Press the locking levers down ↓ to lock the casters and up ↑ to unlock them.

⚠️ **CAUTION**

Always lock the brakes ↓ before setting the X-ray tube unit.

**Overhead (ceiling-mounted) model with SIROLUX FANTASTIC**
Centro Column with HELIODENT DS wall model

Kit:
Adapter for HELIODENT DS X-ray unit on Centro Column of KaVo company, with long support arm (1810mm) and HELIODENT DS wall model only, separate control electronics with timer and release.

Split Concept with HELIODENT DS wall model

Split Concept adapter only for long support arm (1810mm) with x-ray tube and HELIODENT DS wall model only, separate control electronics with timer and release.
## 11 Exposure times

### 11.1 Possible exposure times

<table>
<thead>
<tr>
<th>Seconds</th>
<th>0.01</th>
<th>0.02</th>
<th>0.03</th>
<th>0.04</th>
<th>0.05</th>
<th>0.06</th>
<th>0.08</th>
<th>0.10</th>
<th>0.12</th>
<th>0.16</th>
<th>0.20</th>
<th>0.25</th>
<th>0.32</th>
<th>0.40</th>
<th>0.50</th>
<th>0.64</th>
<th>0.80</th>
<th>1.00</th>
<th>1.25</th>
<th>1.60</th>
<th>2.00</th>
<th>2.50</th>
<th>3.20</th>
<th>3.20</th>
</tr>
</thead>
</table>

### 11.2 Exposure times with 8" source-skin distance tube - Class E film

#### Supramaxillary
- Seconds: 0.08, 0.10, 0.12, 0.16, 0.20, 0.25, 0.32, 0.40, 0.50, 0.64

#### Submaxillary
- Seconds: 0.08, 0.10, 0.12, 0.16, 0.20, 0.25, 0.32, 0.40, 0.50, 0.64

### 11.3 Exposure times with 12" source-skin distance tube - Class E film (Round or square cone)

#### Supramaxillary
- Seconds: 0.20, 0.25, 0.32, 0.40, 0.50, 0.64, 0.80, 1.00

#### Submaxillary
- Seconds: 0.20, 0.25, 0.32, 0.40, 0.50, 0.64, 0.80, 1.00

**NOTE:**
- Set one to two steps **lower** when using class F film types.
- Set three steps **higher** when using Class D film types. **Example:** select 0.40 s instead of 0.20 s. Set 1-2 steps **higher** when using a film holder.
11.4 **Exposure times for Sirona Full Size / Universal X-ray sensors with 8"FHA tube**

The recommended exposure times are limited to the following values from the range of possible exposure times:

<table>
<thead>
<tr>
<th>Tooth Region</th>
<th>Exposure Time</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mandibular front tooth, canine tooth</td>
<td>0.04 s</td>
</tr>
<tr>
<td>Maxillary front tooth</td>
<td>0.05 s</td>
</tr>
<tr>
<td>Mandibular molar</td>
<td>0.06 s</td>
</tr>
<tr>
<td>Maxillary canine tooth, occlusal exposure of molars</td>
<td>0.08 s</td>
</tr>
</tbody>
</table>

**ATTENTION:** The two tables below give only examples for the X-ray sensor exposure times. Please refer to the operating instructions for the X-ray sensors for the correct exposure times.

11.5 **Exposure times for Sirona Full Size / Universal X-ray sensors with 12"FHA tube (Round or square cone)**

The recommended exposure times are limited to the following values from the range of possible exposure times:

<table>
<thead>
<tr>
<th>Tooth Region</th>
<th>Exposure Time</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mandibular front tooth, canine tooth</td>
<td>0.08 s</td>
</tr>
<tr>
<td>Maxillary front tooth</td>
<td>0.10 s</td>
</tr>
<tr>
<td>Mandibular molar</td>
<td>0.12 s</td>
</tr>
<tr>
<td>Maxillary canine tooth, occlusal exposure of molars</td>
<td>0.16 s</td>
</tr>
<tr>
<td>Mandibular molar</td>
<td>0.08 s</td>
</tr>
<tr>
<td>Maxillary canine tooth, occlusal exposure of molars</td>
<td>0.10 s</td>
</tr>
</tbody>
</table>
## Exposure times using tube with 8” SSD - General X-ray sensors

<table>
<thead>
<tr>
<th>Seconds</th>
<th>0.01</th>
<th>0.02</th>
<th>0.03</th>
<th>0.04</th>
<th>0.05</th>
<th>0.06</th>
<th>0.08</th>
<th>0.10</th>
<th>0.12</th>
<th>0.16</th>
<th>0.20</th>
<th>0.25</th>
<th>0.32</th>
</tr>
</thead>
</table>

*These parameters enable an especially high dose reduction for children resulting in increased image noise. If you perceive this noise to be disturbing, we recommend manually increasing the exposure time 1 - 2 steps.

## Exposure times using tube with 12” SSD - General X-ray sensors (Round or square cone)

<table>
<thead>
<tr>
<th>Seconds</th>
<th>0.04</th>
<th>0.05</th>
<th>0.06</th>
<th>0.08</th>
<th>0.10</th>
<th>0.12</th>
<th>0.16</th>
<th>0.20</th>
<th>0.25</th>
<th>0.32</th>
</tr>
</thead>
</table>

*NOTE: When altering the basic setting (see Section 5.4) a new SIDEXIS system setting is required. Please contact your nearest service technician.*
12 Care of surfaces

Cleaning

Remove dirt, grime and disinfectant residue with mild, household detergent. Do not allow fluids to enter the ventilation slits!

Any medicinal agents accidentally spilled onto surfaces should be wiped off immediately.

Disinfecting

Disinfect using either spray-on or wipe-on surface disinfectants. Be sure to observe all instructions and restrictions specified by the manufacturer. Use only tested and approved agents!

Never use: agents containing active ingredients or additives such as carbolic acid, peracetic acid, peroxide or other oxygen-cleaving agents, sodium hypochlorite or any iodine-cleaving agents.

Caring for the film holder

Please observe the hygiene instructions of the film holder manufacturer.

Caring for the SiDEXIS sensor holder system

Please observe the disinfection and sterilization instructions of DENTSPLY RINN.
Errors occurring during the self-test are indicated by a numeric error code of 1 - 4.

<table>
<thead>
<tr>
<th>Error number</th>
<th>Description of cause</th>
<th>Corrective action</th>
</tr>
</thead>
<tbody>
<tr>
<td>No. 1</td>
<td>The cable connecting the X-ray tube unit to the generator may be interrupted (kV line).</td>
<td>Switch unit OFF. Contact your service technician.</td>
</tr>
<tr>
<td>No. 2</td>
<td>X-ray was not exposed (mA/kV_{max} error).</td>
<td>Switch unit OFF. Contact your service technician.</td>
</tr>
<tr>
<td>No. 3</td>
<td>D button was pressed during switch-on phase.</td>
<td>Switch unit OFF and ON again. Be sure not to press the D button while doing so.</td>
</tr>
<tr>
<td>No. 4</td>
<td>Release button was pressed during switch-on phase.</td>
<td>Switch unit OFF and ON again. Be sure not to press the release button while doing so.</td>
</tr>
</tbody>
</table>

**CAUTION**

*If an error is displayed again after switching the unit off and then on again, please contact your service technician.*
14 Inspection and maintenance

Inspection and maintenance work must be performed at regular intervals to protect the safety and health of patients, users and third parties.

14.1 Annual inspection performed by the operator or other authorized personnel

As the operator, you should ensure the safety and reliability of your system by performing maintenance on it at regular intervals (at least once annually) or having this work performed by your dental dealership.

The information in the document 'Inspection and maintenance' supplied will be of help to you here.

14.2 Maintenance performed by the service technician

In addition to the scheduled annual inspection by the user or persons contracted to perform this, a maintenance inspection must be performed after 4, 7, 10 and then every two years.

The information in the document 'Inspection and maintenance' supplied will be of help to the service technician here.

14.3 Checking image quality

At regular intervals, however at least once a year, the user must evaluate the image quality.

With the use of digital image receptors, the increase in image post-processing effort making use of the brightness or contrast control (e.g. for SIDEXIS) can be taken as an evaluation criterion.

For X-rays requiring developing of films, an increase in the exposure time serves as the evaluation criterion.

If these evaluation criteria are fulfilled independently of the patient anatomy and of possible sources of error, such as developing of film or patient positioning, contact a service technician immediately in order to eliminate possible unit faults.

In addition it is necessary to observe country-specific requirements.

Furthermore, we would like to call your attention to the brochures of film manufacturers and also to our Quality Image Service. For this, please contact your dental dealer or the manufacturer directly.