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1. System Description

INTRABEAM® is a mobile radiation instrument which received both FDA and CE approval for the irradiation of all solid tumors in 1999. Treatment can be performed in operating rooms, special radiation protection measures are normally not required. The mobile radiation source of INTRABEAM® produces low-energy x-ray radiation which is emitted isotropically (equally distributed). The INTRABEAM® carrier system with six degrees of freedom, weight compensation and magnetic brakes ensures easy, flexible and precise positioning of the source into the targeted area. Ideally integrated in the INTRABEAM® Cart, the control unit ensures exact setting and monitoring of the desired dose. INTRABEAM® is a mobile system, which can be used parallel in multiple operating rooms.
2. Mobility of the System & Interaction of the System Components

All INTRABEAM® components are completely flexible and movable. Most components, such as the 1.6 kg light x-ray source, are stored in a storage container inside the INTRABEAM® Cart. The cart and floor stand can be easily stowed away with all components when not in use. With its open space on top of the cart, the system quality check can be done easily on the cart. The touch pad terminal, control console, dosimeter and everything else needed for system quality assurance and the treatment are mounted ergonomically on the cart. Large casters and guiding rollers ensure easy transportation inside and outside the operating room. The set up of the cart also ensures quick and easy cleaning in the OR. The cart and floor stand can be moved through normal doors and through elevators if necessary. When in use, the x-ray source is inserted in the arm of the INTRABEAM® carrier. With 12 casters in the base, the carrier can be moved smoothly to any position in the OR. Weight compensation and six axes provide enough freedom to bring the x-ray source in every position needed for treatment. Electromagnetic brakes hold the source into the exact position during treatment. The operator can control the system at any time from the control panel at the INTRABEAM® Cart during the process.
3. Dimensions of the System Components

**INTRABEAM® X-Ray Source 4**

Weight: 1.6 kg / 3.57 lbs  
Dimensions: 70 x 175 x 110 mm (width x height x length)

**INTRABEAM® Cart**

Weight of cart unloaded, incl. permanently mounted user terminal: approx. 105 kg / 231 lbs  
Payload max.: 95 kg / 209.43 lbs  
Dimensions: 900 x 1690 x 600 mm / 35.43” x 66.53” x 23.62”

**INTRABEAM® Floor Stand**

Weight: 275 kg / 606 lbs  
Transport Position: 740 x 1940 x 1500 mm / 29.13” x 76.38” x 59.06”

**Consumables:**

**INTRABEAM® Radiation Shield Flat 20 cm x 20 cm**

(10 per box)  
Material: Tungsten

**INTRABEAM® Drapes**

(5 per box)
Floor stand
Dimensional drawings
1 Working position (example)
2 Transport position
4. X-ray Source and Beam Characteristics

The mobile radiation source of INTRABEAM® accelerates electrons through the 100 mm drift tube with a maximum voltage of 50 kV onto a gold target where the low-energy x-ray radiation is generated and then emitted isotropically.

4.1 Beam energies

40 kV / 10, 20 or 40 μA
50 kV / 5, 10, 20 or 40 μA

Spherical dose distribution of the x-rays. The steep drop in the dosage ensures that most of the radiation remains in the targeted tissue.
4.2 Beam characteristics

<table>
<thead>
<tr>
<th>Diameter of the target area [mm]</th>
<th>Approx. Radiation intensity in the periphery [Gy/Minute]</th>
<th>Approx. Treatment time @ 15 Gy [Minute]</th>
</tr>
</thead>
<tbody>
<tr>
<td>10</td>
<td>9.8</td>
<td>1.5</td>
</tr>
<tr>
<td>20</td>
<td>2.3</td>
<td>6.5</td>
</tr>
<tr>
<td>30</td>
<td>0.9</td>
<td>16.7</td>
</tr>
<tr>
<td>40</td>
<td>0.4</td>
<td>37.5</td>
</tr>
<tr>
<td>50</td>
<td>0.25</td>
<td>60</td>
</tr>
</tbody>
</table>

All displayed values are valid for the operation of the XRS4 at 50 kV and 40 μA.

4.3 Beam allocation

- Point-source type emission
- Dose isotropic around XRS 4 isocenter
- Sharp dose fall off (approx. 1/r³) in soft tissue or equivalent
- Positional uncertainty of delivered dose less ± 1 mm at 40 mm treatment diameter.

4.4 On-site dose monitoring

An internal radiation monitor (IRM) detects the back bouncing counts and assures on-site monitoring. The IRM result is displayed on the treatment screen of the control terminal so that the operator knows which dose is being delivered at any time during treatment.
5. Dosimetry System

A full set of quality assurance and dosimetry tools are provided with the INTRABEAM®. The factory-calibrated system is delivered with the specific depth dose curves and a reference measurement with the ion chamber belonging to the system. Every day, a two-step quality control check ensures that all parameters such as isotropy, internal radiation monitor and output work do not exceed the tolerances defined during calibration. For commissioning, a completely shielded, manual adjustable water phantom can be used to remeasure the depth dose curve.

5.1 Consignment Sale
(System calibration by Carl Zeiss and with the water phantom)

A water phantom with a highly precise movement technique enables the physicist to position the tip of the XRS4 exactly above or beside the ion chambers inside the water. Accurate positioning and stepping of the source ensures the remeasurement of the depth dose curve. Even the measurement of a depth dose curve is possible.

5.2 QA Check (QA Tools)

Inside the so called PDA, five diodes in each direction measure the radiation of the XRS4. The objective of this test is to ensure the sphericity of the beam. With the PAICH the output in the z-axis can be checked. An ion chamber is mounted in the probe adjuster in a way that the ion chamber window is right above the tip of the XRS4. In this test, the internal radiation monitor and the output are checked.
5.3 Online control (IRM)

Prior to each treatment session, the x-ray source must be subjected to a verification process comprising at least two levels before it is enabled by the software for radiation emission.

- Verification of the isotropy of the radiation field. A photodiode array (PDA) is used to test the radiation field in all 5 axes. Only if all diodes measure identical radiation, i.e. if the field is spherical within specified tolerances, will the system be released for the next verification step.
- Verification of the radiation output using an external ionization chamber is only possible after successful completion of the PDA test. The counts measured by the internal radiation monitor are compared with the reading of the ionization chamber. The x-ray source is not enabled for treatment planning until a coefficient has been computed.

5.4 Radiation protection during dosimetry

Every tool provided with the INTRABEAM® is completely self shielded and does not require any additional radiation protection.
6. Safety Concept (Interlocks)

A closed, meshed interlock system controls the INTRABEAM® during treatment and quality assurance. It prevents the user against e.g. unintended radiation emissions, unintended access to the controlled area and incorrect radiation data transmission (see 6.1 – 6.3). This safety concept is also responsible for ensuring the proper work during treatment and will adjust and sound an alert if anything else occurs. To obtain the license for operation of the INTRABEAM®, it may be necessary to create a safety strategy and/or failure analysis, a hospital radiation protection manual and an emergency plan. Suggestions on how to establish this are provided by Carl Zeiss.

6.1 Interlock to prevent unintended radiation emission

Unintended radiation emissions are prevented by an optical interlock system and a multilayered inquiry in the software.

- The optical interlock detects whether the appropriate verification device has been correctly attached to the x-ray source. Only then and only when the device protects the environment from radiation, will the x-ray source be enabled for radiation emission. If the device is removed during the test, radiation emission is immediately deactivated by the interlock.

- The user is requested via the user interface to press the start button at least twice to confirm that he really wants to perform the test. The unintended execution of a verification test is not possible.

- The optical interlock enables the system to detect if the x-ray source has been correctly mounted on the stereotactic frame adapter or the INTRABEAM® Floor Stand. The system also detects whether the applicator has been correctly attached to the INTRABEAM® Floor Stand. The x-ray source will only be enabled for radiation emission if it has been correctly mounted on the carrier system intended for it.

- Once the x-ray source has been successfully connected to the carrier system, the user must follow the menu prompts on the user interface to actively set the system status to ready for radiation emission. Radiation will only be emitted if the user presses the Start button again.

- As soon as the system is ready for radiation emission, this ready status is indicated by an acoustic signal. This warns the user that radiation will be emitted when the Start button is pressed again.

- Radiation emission can only be started by trained and authorized staff after the necessary verification steps have been completed, the dose has been confirmed by a second person verified by a password and the system has been set to the treatment mode.
6.2 Interlock to prevent unintended access to the controlled area

Protection against unintended access to the controlled area
- Radiation emission is acoustically and visually indicated on the x-ray source and the user interface. Persons entering the controlled area are able to obtain information on the radiation status of the system.
- In addition, further safety systems such as an external warning lamp or a door contact switch can be connected with the INTRABEAM® via an external interlock switch. If, for example, the external interlock is activated by opening of the door during a treatment session, radiation emission is instantly interrupted automatically and can only be resumed after the interlock has been closed and continued radiation emission has been confirmed.

6.3 Interlock to prevent incorrect radiation data transmission

Protection against incorrect radiation data transmission
- Every signal transmitted by the control console to the x-ray source is returned by the x-ray source and checked for completeness and correctness by the control console. If the signal is not correctly returned by the x-ray source, radiation emission is stopped or not started.
- The count rate is constantly monitored during treatment by the internal radiation monitor. If the count rate deviates from the planned rate by more than 10%, radiation emission is stopped.
- The ratio between the planned treatment time and the count rate is constantly monitored. In the event of any deviations, radiation emission is stopped.

6.4 Interlock to prevent incorrect dose entry

- The dose to be administered must only be prescribed by an appropriately trained, authorized doctor (this is a legal requirement in most countries; information during system training and note in the user manual).
- The dose is entered by one person (usually a physicist), and a second person, who must be a doctor (this is checked via the profile of the user name), must verify the dose planning and confirm it by entering a password.
6.5 Interlock to prevent interference with the application software

• The INTRABEAM is a closed system. Only data conforming to the specific data format of the application software can be loaded via CD/DVD.

6.6 Electrical safety

• The system has been successfully subjected to the conformity assessment procedure and complies with the essential requirements stipulated in Council Directive 93/42/EEC and with IEC 60601-1, EN 60601-1, UL 60601-1, CAN/CSA-C22.2 No.601.1, IEC 60601-1-2 and IEC 60601-2-8.
7. Application Options

The physical and radiobiological effect of the low energy x-rays produced from the XRS4 INTRABEAM® is used in nearly the whole body side (except heart and central circulation system). Various types of different applicators suitable for the different cavities are used to fill out the tumor bed after removal of the tumor. Carl Zeiss continuously works on the development of new applicator types for additional treatment applicators.

7.1 INTRABEAM® Spherical Applicator

Spherical applicators are available with diameters of 1.5 to 5 cm in half cm increments. The applicator fills out the entire wound cavity after tumor removal. The tumor bed tissue is adapted smoothly around the applicator surface. The x-ray probe is located then in the center of the applicator and therefore also of the tumor cavity. The material of the applicator is medical grade acrylic and can be steam sterilized.
7.2 INTRABEAM® Balloon Applicator *

Balloon applicators are restricted to breast cancer therapy and are available with diameters of 3 to 5 cm in half cm increments. They can be inserted through a small channel into the tumor bed. During the time the balloon is inflated the solid spherical shape does not change. After treatment the balloon can be deflated and removed through the channel without the need for another operation. The balloon applicator comes in a sterile set with accessories to place and remove the applicator.

7.3 INTRABEAM® Needle Applicator **

To perform tumor bed treatment in lesions which can be reached only through very small incisions the needle applicator can be used.

* The INTRABEAM® Balloon Applicator is FDA approved. The INTRABEAM® Balloon Applicator is pending CE clearance.

** The INTRABEAM® Needle Applicator is pending FDA clearance. The INTRABEAM® Needle Applicator is CE approved.
8. Technical Requirements

8.1 Electrical requirements

Two outlets are necessary to use the INTRABEAM® during treatment – one for the floor stand and one for all components on the cart which are connected together with a medical insulating transformer.

INTRABEAM® Floor Stand
Rated voltage 100V / 115V/ 230V
Power consumption max. 400 VA
Rated frequency 50 - 60 Hz
Electrical standard; Complies with IEC 60601-1 / EN 60601-1; UL 60601-1;
Approval CAN/CSA-C22.2 No. 601.1-M90
Protection class I Degree of protection IPX0
Type B equipment
Product classification I in acc. with 93/42/EEC Annex IX

INTRABEAM® Cart Medical Insulating Transformer
Rated voltage 100V - 240 V
Rated frequency 50 / 60 Hz
Power input 600 VA
Power output: 230 V
Case protection IP 20
Protection Class I

8.2 Ambient requirements*

Operation:
Temperature +15 °C .... +40 °C
59 °F.....104°F
Relative humidity: 30%....75%
Air pressure: 700 hPa.....1060 hPa

Transportation and Storage:
Temperature: -20 °C .... +70 °C
-4 °F....158 °F
Relative humidity: 10%.....90% (without condensation)
Air pressure: 500 hPa.....1060 hPa

*Requirements could differ for accessories. Please see separate user manuals.
9. Processing (Cleaning/Disinfection/Sterilization)

All non-sterile components of the INTRABEAM® can be easily cleaned and disinfected. Large casters and nearly no tight cleavings ensure simple and quick cleaning of the cart. The floor stand can also be cleaned in a very short time.

<table>
<thead>
<tr>
<th>Component</th>
<th>Cleaning</th>
<th>Disinfection</th>
<th>Single Use/ Sterilizable</th>
</tr>
</thead>
<tbody>
<tr>
<td>Floor Stand</td>
<td>c</td>
<td>b</td>
<td>–</td>
</tr>
<tr>
<td>INTRABEAM® Cart</td>
<td>c</td>
<td>b</td>
<td>–</td>
</tr>
<tr>
<td>Control Console PRS 500</td>
<td>c</td>
<td>a</td>
<td>–</td>
</tr>
<tr>
<td>User Terminal</td>
<td>c</td>
<td>b</td>
<td>–</td>
</tr>
<tr>
<td>Keyboard</td>
<td>c</td>
<td>a</td>
<td>–</td>
</tr>
<tr>
<td>X-ray Source 4</td>
<td>c</td>
<td>a</td>
<td>ETO</td>
</tr>
<tr>
<td>Quality Tools (PDA/PAICH)</td>
<td>c</td>
<td>a</td>
<td>–</td>
</tr>
<tr>
<td>Cables (XRS/QA)</td>
<td>c</td>
<td>a</td>
<td>–</td>
</tr>
<tr>
<td>V-/X- Block</td>
<td>c</td>
<td>a</td>
<td>–</td>
</tr>
<tr>
<td>INTRABEAM® Water Phantom</td>
<td>d</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Transportation Trays (XRS4/QA)</td>
<td>c</td>
<td>a</td>
<td>–</td>
</tr>
<tr>
<td>INTRABEAM® Spherical Applicator</td>
<td>look on page 19</td>
<td>X</td>
<td>sterilizable - steam</td>
</tr>
<tr>
<td>INTRABEAM® Balloon Applicator</td>
<td>–</td>
<td>–</td>
<td>single use - sterile</td>
</tr>
<tr>
<td>INTRABEAM® Needle Applicator</td>
<td>–</td>
<td>–</td>
<td>single use - sterile</td>
</tr>
<tr>
<td>INTRABEAM® Flat Shields</td>
<td>–</td>
<td>–</td>
<td>single use - sterile</td>
</tr>
<tr>
<td>INTRABEAM® Drapes</td>
<td>–</td>
<td>–</td>
<td>single use - sterile</td>
</tr>
</tbody>
</table>

a = 4–7 % Hypochlorid, b = Meliseltol, c = wipe moist, d = ethyl alcohol, distilled water (1:1) plus a dash of household dish-washing liquid also mentioned in the detailed user manuals.
The processing method of the INTRABEAM® Spherical Applicators was validated by Carl Zeiss. The INTRABEAM® Spherical Applicators can be processed with the following method:

If possible, you should use a machine for cleaning and disinfection. The manual procedure should only be used when a machine is not available, as it is clearly less effective and reproducible.

- Never clean the applicators and sterilization containers with metal brushes or steel wool.
- Never clean the applicators and sterilization containers using ultrasound procedures.
- The applicators and sterilization containers must not be exposed to temperatures higher than 141°C (286°F).
- Prepare cleaning and disinfecting agents that meet the following requirements:
  - Aldehyde-free disinfecting agent (otherwise blood contamination remains). The disinfecting agent must be certified in accordance with VAH/DGHM or FDA approval or CE label stipulations.
  - The disinfecting agent is compatible with the cleaning agent used.
  - The disinfecting agent is generally suitable for cleaning and disinfecting devices made of metal and plastic.
  - The disinfecting agent must not contain the following:
    - organic, mineral and oxidizing acids (minimum permissible pH value: 5.5)
    - strong bases (maximum permissible pH value: 8.5, recommended: neutral, enzymatic cleaning agent)
    - organic solvents (e.g. alcohol, ether, ketone, benzine)
    - oxidants (e.g. hydrogen peroxide)
    - halogens (chlorine, iodine, bromine)
    - aromatic/halogenated hydrocarbons
    - Rinsing with sterile or low-germ (maximum 10 microbes/ml) and lowendotoxin (maximum 0.25 endotoxin units/ml) water (e.g. purified water/highly purified water)
- Only use the steam sterilization method (fractioned vacuum method) described in this manual for sterilization of the cleaned and disinfected applicators. Other procedures are not approved.
- Do not use gravitation methods, hot air sterilization, radiation sterilization, formaldehyde sterilization, ethylene oxide sterilization or plasma sterilization.
- For steam sterilization, please take the following factors into account:
  - The steam sterilization method is validated according to DIN EN ISO 17665 (previously: DIN EN 554/ANSI AAMI ISO 11134) (valid IQ/OQ (commissioning) and product-specific performance assessment (PQ))
  - The maximum sterilization temperature is 138°C (280°F; plus tolerance according to DIN EN ISO 17665 (previously: DIN EN 554/ ANSI AAMI ISO 11134))
  - With the fractioned vacuum method, the sterilization time (exposure time at sterilization temperature) is at least 5 min1 at 132°C (270°F)/ 134°C (273°F) 1or 18 min (prion inactivation)

Please see the user manual for detailed information on cleaning, disinfection and sterilization of the INTRABEAM® Spherical Applicators.