Instructions for Use: The Pink Pad®

BEFORE USING PRODUCT, READ THE FOLLOWING INFORMATION THOROUGHLY.

DESCRIPTION

The Pink Pad is a single use system for use in surgical procedures. The system consists of a proprietary formulation for the pink foam pad, non-woven lift sheet, body straps, head rests, and boot liners.

INDICATIONS

The Pink Pad is a single use, non-sterile device to be used during surgery to provide patient protection from injury.

INTENDED PURPOSE AND USER

This device is intended for use by trained healthcare professionals only.

CONTRAINDICATIONS

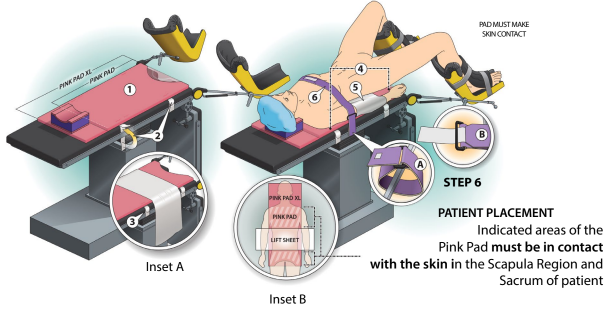
This device is not designed, sold, or intended for use except as indicated.

KNOWLEDGE AND USE

Professional use requires knowledge of this instruction for use. Device use limited to surgical operating room in a hospital or surgery center.

CLINICAL BENEFITS

To aid hospital facilities in providing a safe and effective method of management of the patient for pressure ulcers and non-movement for patients in the Trendelenburg position.



PREPARATION AND USE

- Place The Pink Pad at the very edge of the middle table segment nearest the perineal cutout, positioning the white hook & loop straps on the underside of The Pink Pad in direct contact with the table pad's surface. IMPORTANT: DO NOT use any table covers, linens, other transfer devices or materials between The Pink[®] Pad and surface of the surgical table. The Pink Pad must contact the surface of the table pad. The "This Side Up" tags should be facing upward.
- 2. Attach the white hook & loop straps of The Pink Pad to the surgical bed rails by looping under the rail, as shown, and affixing the ends of the hook & loop to each other.
- Lay the lift sheet over the pad, centered between the hook & loop straps as shown (inset A). The lift sheet should cover only the portion of the pad that will be addressing the small of the patient's back - below the Scapula Region and above the Sacrum.
- 4. Follow hospital protocol for intubation. Then properly position patient on pad. IMPORTANT THE PATIENT'S SKIN MUST MAKE DIRECT CONTACT WITH THE PINK PAD. The skin of the Scapula Region and Sacrum must contact the pad surface (inset B). Utilize the included lift sheet to carefully lift the patient up and off the pad to reposition as needed for safe and proper application of stirrups. Do not drag patient on pad. Make sure pad remains completely flat at all times.

- 5. Tuck arms as shown or per hospital protocol.
- Attach the body strap as follows:
 - A. Place the strap component with the small hook & loop square around the table's accessory rail and through strap buckle
 - B. Repeat the above step on the other side of the table with the remaining strap component with the hook & loop's hook side facing downward. Join straps together.

DISPOSAL

After use, the Pink Pad should be disposed of in accordance with hospital policy.

WARNINGS

- This device was designed, test and manufactured for single patient use only. Reuse or reprocessing of this device may lead to its failure or subsequent injury.
- Reprocessing of this device may create the risk of contamination and patient infection.
- Do not reuse or reprocess this device.
- After use, the Pink Pad should be disposed of in accordance with hospital policy.
- The Pink Pad should be stored in a clean, dry location at room temperature prior to use. Avoid prolonged exposure to elevated temperatures.

PRECAUTIONS

- The Pink Pad should be used in accordance with the instructions for use and any contraindications, warnings or precautions provided by the manufacturer of the associated instrument.
- Before using The Pink Pad, ensure that the O.R. table pad is securely affixed to the O.R. table and is clean and free of residue.
- Be sure to follow your facility's policies and guidelines for frequency of patient monitoring. Check skin for integrity and proper circulation. Product is to be used by licensed medical professional only.
- Care should be taken to safeguard The Pink Pad from exposure to prep solutions.
- Notice to the User and/or patient that any serious incident that has occurred in relation to the device should be reported and the competent authority of the Member State in which the user and/or patient is established, as well as, Xodus Medical and its Authorized Representative.

TECHNICAL SPECIFICATIONS

- Materials of manufacture include:
 - » Polyurethane foam, synthetic adhesive, nylon strap
- Shelf Life indefinite

Non-steril

STORAGE, TRANSPORT, AND OPERATIONAL CONDITIONS

- The Pink Pad should be stored in a clean, dry location at room temperature prior to use. Avoid prolonged exposure to elevated temperatures.
- During ALL handling and storage, assure that the pad is flat. Do not roll or fold the pad.







CE n/eifu Consult instructions for use or consult electronic instructions for use

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Accompanying Documents

LATEX Not Made with Natural Rubber Latex



Do Not Use If The

Package Is Damaged

Or Opened



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FDA REGISTERED ISO 13485 CERTIFIED





