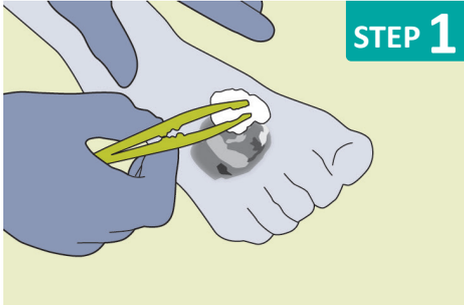


Silicone Border (Instructions For Use)

Asguard Silicone Border dressing (Type I) improves absorption and retention by the combination of a foam and superabsorbent layer. It absorbs exudate through silicone perforations from the wound and provides a moist wound healing environment to promote wound healing. The wound pad is partly perforated with cut to improve the product extensibility. The silicone wound contact layer reduces pain and trauma during dressing changes, and enhances patient's comfort. The waterproof outer layer protects the wound from dirt and bacteria.



STEP 1

Clean the wound area in accordance with normal procedures. Ensure the peri-wound skin is dry.



STEP 2

Select a dressing size of Asguard Silicone Border large enough to overlap the wound edges by at least 1-2 cm for smaller sizes (sizes up to 12.5x12.5cm), and 3-4cm for larger sizes in order to protect the surrounding skin from maceration and fix the dressing securely.



STEP 3

Remove the release film and apply the adherent side to the wound. Smooth the dressing from the center to the edge without leaving any gap, press the dressing edge to ensure good adhesion.



STEP 4

To remove the dressing: gently lift the corner of the dressing and slowly peel.

NOTE

1. The dressing change interval may be several days. Change the dressing before it is fully saturated, at signs of leakage, or as indicated by clinical practice. The dressing may be left in place up to 7 days.
2. When necessary, fixate the dressing with a bandage or other fixation.

STORAGE & TRANSPORT

The product should be stored in dry conditions.

Keep the product away from direct sunlight and keep dry.

DISPOSAL

Disposal should be handled according to local environment procedures.

At Sentry, we are dedicated to upholding sustainable principles to safeguard our planet, people, and future, and we strive to consistently act responsibly in doing so.

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Silicone Border (Instructions For Use)

FUNCTION PRINCIPLE

Asguard Silicone Border dressing absorbs exudate and provides a moist wound healing environment to promote wound healing. The perforated silicone contact layer provides gentle adhesion and secure fixation and does not stick to the moist wound resulting in less pain for patients. The polyurethane foam layer absorbs exudates and transfers it to a superabsorbent layer using a non-woven spreading layer. Super absorbent fibres absorb and retain exudate by forming a gel, minimizing the risk for maceration. The waterproof outer layer protects the wound from dirt and bacteria.

INTENDED PURPOSE & INDICATIONS

INTENDED PURPOSE

Asguard Silicone Border dressing is a wound dressing that absorbs wound exudate and provides a moist wound healing environment to promote wound healing. It can also prevent pressure ulcer.

INDICATIONS

Asguard Silicone Border dressing is designed for a wide range of exuding wounds such as pressure ulcers, leg and foot ulcers, traumatic wounds (e.g. skin tears) and surgical wounds. It can also be used on dry/necrotic wounds in combination with gels. The dressing can reduce postoperative blistering, and may also be used as part of a prophylactic therapy to help prevent skin damage, e.g. pressure ulcers.

INTENDED USER

The product shall be used by or under the supervision of a healthcare professional.

INTENDED PATIENT POPULATION

Patient who has the indications: pressure ulcers, leg and foot ulcers, traumatic wounds (e.g. skin tears), surgical wounds and dry/necrotic wounds.

CLINICAL BENEFIT

1. As an adjunct to pressure ulcer prevention: reduce the incidence of pressure ulcers when combined with usual pressure injury prevention practices;
2. For wound management: reduce patient pain at dressing change/removal.
3. For wound management: promote wound healing.

CONTRAINDICATIONS

1. Check the wound for signs of infection before use, if infection occurs, see a health care professional.
2. Do not use on the patients with a known hypersensitivity to the product itself or to its components.
3. Do not use on third degree burns.

COMPLICATIONS OR SIDE-EFFECTS

Very occasionally, there will be some potential risks associated with the use of Asguard Silicone Border Dressing with the following:

- Adverse reactions (eczema, erythema, pruritus, blister formation, skin rash, allergy etc.)
- Wound infections
- Risk of maceration (fluid leakage)
- Risk of skin stripping

WARNINGS

1. Do not reuse. Reuse will cause cross-contamination.
2. Do not use if package is damaged or open.
3. Do not use together with oxidizing agents such as hypochlorite solutions or hydrogen peroxide.
4. Due to the gentle adhesive used, do not use this dressing to secure other medical devices to the patient.
5. If reddening or sensitization occurs discontinue use and see a healthcare professional.

OTHER INFORMATION

1. The foam may change colour to more yellow when exposed to light, air and/or heat. This has no influence on product properties.
2. The intended user does not need training for this product.
3. If have any serious incident that has occurred in relation to the device, please report to the manufacturer and the competent authority.
4. The specified goods should only be used under the supervision of a healthcare professional.
5. If you are concerned about your wound, consult a healthcare professional.

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POUCH - PAPER



POUCH - FILM



BOX

