**3M™ Tegaderm™ Hydrogel**

**Hydrogel Wound Filler**

**Description:**
3M™ Tegaderm™ Hydrogel Wound Filler is a sterile, non-preserved, amorphous hydrogel formulated to provide moisture to a dry wound, maintain a moist wound environment which enhances wound healing and helps prevent wound desiccation, and promote autolytic debridement of a dry wound by providing moisture to devitalized tissue.

Ingredients: water, propylene glycol, guar gum and sodium tetraborate.

**Indications for use:**
Tegaderm™ Hydrogel Wound Filler is indicated as a primary wound filler for non-to minimally draining partial and full thickness dermal wounds including:

- pressure ulcers
- venous ulcers
- arterial ulcers
- diabetic ulcers
- abrasions and lacerations
- grafts and donor sites
- post operative surgical wounds
- cavity wounds
- wounds exhibiting dry eschar and fibrinous slough

Tegaderm™ Hydrogel Wound Filler may be applied to sterile gauze and used to lightly pack tunneled or undermined wounds. Tegaderm™ Hydrogel Wound Filler may be used on infected wounds only under the care of a health care professional. This product is not designed, sold, or intended for use except as indicated.

**Contraindications:**
None known.

**Warnings:**
None known.

**Precautions:**
1. For external use only.
2. Do not use on patients with a known sensitivity to propylene glycol.
3. Treatment of any wound should be part of a well-defined plan and under the supervision of a health care professional.
4. Bleeding should be controlled before application of wound fillers or cover dressings.
5. When using wound fillers or dressings, the wound may initially appear larger in size and depth as unnecessary tissue is cleaned away. This increase should be accompanied by an improved appearance of the wound. If the wound gets larger after the first few dressing changes, consult a health care professional.
6. Observe the wound for signs of infection. Consult a health care professional if any of the following are noted: fever, increased pain, redness, bleeding, swelling, unusual odor, increased drainage or cloudy or foul drainage.
7. Tegaderm™ Hydrogel Wound Filler may be used on infected wounds only under the care of a health care professional.
8. Rarely, with wound filler or dressing use, irritation (redness), or maceration (whitening) of the surrounding skin, or hypergranulation (excessive tissue formation in the wound) may develop. Should this occur, consult a health care professional.
9. If the wound does not begin to show signs of healing or if any other unexpected symptoms occur, consult a health care professional.
10. Single use only, once opened this product should not be reused due to risk of contamination and infection.
11. Tegaderm™ Hydrogel Wound Filler must not be ingested and must be kept away from children and animals.

**Directions for Use:**

**Note:** Follow facility guidelines for infection control.

1. Cleanse wound and surrounding skin according to facility policy.

   **Note:** If periwound skin is fragile or exposure to wound exudate is likely, apply a barrier film such as 3M™ Cavilon™ No Sting Barrier Film. Allow the barrier film to dry before dressing application.

2. Apply enough Tegaderm™ Hydrogel Wound Filler to cover the wound base and any necrotic tissue, to a depth of approximately 5 mm thick, with no overlap onto the surrounding skin.

3. Alternatively, a sterile gauze pad may be saturated with Tegaderm™ Hydrogel Wound Filler and placed into the wound with no overlap onto the surrounding skin.

4. Apply an appropriate cover dressing to help manage the wound drainage and maintain a moist wound environment.

**Removing the Dressing:**

1. Change the dressing as needed. Tegaderm™ Hydrogel Wound Filler may be left in place for up to 3 days. Frequency of changing will depend on factors such as the type of wound and volume of drainage.

2. Carefully remove the secondary dressing and dispose of according to local procedures and guidelines.

3. Cleanse or gently flush the wound with sterile saline or wound cleanser to remove necrotic debris and any remaining gel.

4. At the time of dressing change, if the dressing is adhered to the wound surface, saturate with wound cleanser or sterile normal saline, allow the dressing to soften, and gently remove.

5. Avoid forceful removal of the dressing to minimize disruption of the wound.

**Storage and Shelf Life:**
Store at temperatures <25°C / 77°F.
Store in a dry and cool place.
For shelf life, refer to the expiration date printed on each box and tube.

**How Supplied:**
Supplied in boxes of 10 individual tubes.
Sterility of product is guaranteed unless the individual tube is damaged or open.
If you have any questions or comments, in the USA please contact the 3M Health Care Customer Helpline at 1-800-228-3957.
For further information outside the United States, contact your local 3M representative or contact us at www.3M.com and select your country.

**Caution Statement:** Federal Law restricts the device to sale by or on the order of a licensed health care professional.

**Explanation of Symbols**

- **Not Made With Natural Rubber Latex**
- **Caution, see instructions for use**
- **Do not use if package is damaged**
- **Do not reuse**
- **Use by date**
- **Batch code**
- **Manufacturer**
- **Date of manufacture**
- **Sterilized using steam**
- **Do not resterilize**
- **Upper limit of temperature**

Made in UK for

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