

Kendall™

AMD Antimicrobial Fenestrated Foam Disc Dressing

Description: Kendall™ AMD Antimicrobial Fenestrated Foam Disc dressings provide an ideal dressing to cover wounds resulting from percutaneous devices including vascular and non-vascular catheters. This highly absorbent, non-linting dressing is designed to provide protection and cushioning at these sites. Kendall™ AMD Antimicrobial Fenestrated Foam Disc dressings are semi-occlusive allowing the exchange of gases such as oxygen and water vapour. The soft flexible nature of the Kendall™ AMD Antimicrobial Fenestrated Foam Disc dressing allows it to conform easily to all body contours. Kendall™ AMD Antimicrobial Fenestrated Foam Disc dressings are non-adherent and non-drying, making dressing changes easy and minimising pain. Kendall™ AMD Antimicrobial Fenestrated Foam Disc dressings contain the antimicrobial agent Polyhexamethylene biguanide hydrochloride (PHMB) and provide an antimicrobial barrier to bacteria penetration through the dressing and prevents colonisation and proliferation of bacteria within the dressing for up to 7 days.

Indications for Use: Kendall™ AMD Antimicrobial Fenestrated Foam Disc dressings are ideal dressings in the local management of exudate that may occur at surgically induced body exit sites such as IV catheters, central venous lines, arterial catheters, dialysis catheters, peripherally inserted coronary catheters, mid-line catheters, tracheostomy, G-tube, J-tube, Penrose drain, chest tube, nephrostomy tube, sump drain, orthopaedic pins and epidural catheters. The antimicrobial activity of the PHMB in the KendallTM AMD Antimicrobial Fenestrated Foam Disc dressings help to resist bacterial colonisation within the foam dressing and inhibit bacterial penetration through the foam dressing.

Precautions:

- For external use only
- KendallTM AMD Antimicrobial Fenestrated Foam Disc dressings can be used in conjunction with prescribed therapies for the treatment of infections. Dressings are not intended as a primary treatment for infections. If clinical signs of infection are present, please consult a physician.
- Do not use as a primary treatment for full 3rd degree burns.
- Do not use on patients with known sensitivity to PHMB.
- Do not use if primary package is damaged
- Do not re-sterilize.
- Single use only

Allergen Potential: Kendall™ AMD Antimicrobial Fenestrated Foam Disc dressings contain 0.5% PHMB. Exposure to high concentrations (≥ 2.5%) of PHMB has been infrequently associated with contact dermatitis. If dermatitis is observed, discontinue use.

Preparing the Wound Site: Cleanse the wound area if necessary using a non-toxic cleansing solution such as Kendall™ wound cleanser or saline. Dry the wounds surrounding areas thoroughly before dressing application.

Applying the Dressing:

- 1. Prepare the site per facility protocol.
- 2. Aseptically remove Kendall™ AMD Antimicrobial Fenestrated Foam Disc dressing from package.
- 3. Place Kendall™ AMD Antimicrobial Fenestrated Foam Disc dressing around the percutaneous device. The disc is non-oriented and can be applied with either
- 4. For easiest removal during dressing changes, align disc slit under the existing catheter (covering the perforation/slit with the catheter).
- **5.** Approximate perforation/slit edges for best efficacy.
- 6. Secure Kendall™ AMD Antimicrobial Fenestrated Foam Disc dressing and percutaneous device with Kendall™ transparent dressing or other suitable securement dressing such as Kerlix™ AMD antimicrobial bandage roll dressing.
- 7. Change per facility protocol and as needed for excessive drainage. The KendallTM AMD Antimicrobial Fenestrated Foam Disc dressing is effective for up to 7

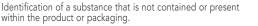
Frequency of Change: The high absorbency and sustained antimicrobial efficacy of KendallTM AMD Antimicrobial Fenestrated Foam Disc dressings allow the user to comply with established treatment protocols for foam dressings, up to one week between dressing changes. In typical use, the frequency of changes will depend upon the nature and condition of the wound and the amount of exudate.

Dressing Change and Removal: The Kendall™ AMD Antimicrobial Fenestrated Foam Disc dressing should be changed when signs of saturation are visible along the edge of the dressing or whenever good nursing practice dictates. To change the dressing, remove the securing bandages or tape and carefully lift Kendall™ AMD Antimicrobial Fenestrated Foam Disc dressing off the insertion site. Kendall™ AMD Antimicrobial Fenestrated Foam Disc dressings are non-adherent, non-adhesive, and non-gelling.

Dressing Expansion: KendallTM AMD Antimicrobial Fenestrated Foam Disc dressings absorb exudate and will expand and 'grow' in size due to the cellular structure of the dressing.

STERILE





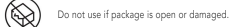




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REV 11/2011

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within the product or packaging.