

## **Australian Government**

# **Department of Health and Aged Care**

Therapeutic Goods Administration

### **Public Summary**

Summary for ARTG Entry: 151997 Coloplast Pty Ltd - Biatain Ibu Non-Adhesive Foam Dressing - Dressing, high-absorbent,

foam/sheet/liquid/powder, non-hydrophilic gel-forming

ARTG entry for Medical Device Included Class III

Sponsor Coloplast Pty Ltd

Postal Address Suite 1 Level 7, 1 Peters Avenue, Mulgrave, VIC, 3170

Australia

ARTG Start Date 30/04/2008

Product Category Medical Device Class III

Status Active

Approval Area Medical Devices

### Conditions

- The inclusion of the kind of device in the ARTG is subject to compliance with all conditions placed or imposed on the ARTG entry. Refer Part 4-5, Division 2 (Conditions) of the Therapeutic Goods (Medical Devices) Regulations 2002 for relevant information.

- Breaching conditions of the inclusion related to the device of the kind may lead to suspension or cancellation of the ARTG entry; may be a criminal offence; and civil penalties may apply.

#### Manufacturers

 Name
 Address

 Coloplast AS
 Holtedam 1

, Humlebaek, 3050

Denmark

### **Products**

## 1. Biatain Ibu Non-Adhesive Foam Dressing - Exudate-absorbent dressing, non-gel

Product Type Single Device Product Effective Date 14/07/2023

**GMDN** 44970 Exudate-absorbent dressing, non-gel

Functional The dressing is a sterile, single use, polyurethane foam dressing which contains ibuprofen (0.5 mg/cm²) homogeneously dispersed throughout the foam. Ibuprofen is released into the wound bed when in contact with wound exudate and may be left in place for up

to 7 days. The dressing should not be changed more than twice daily corresponding to a maximum daily use of 2400 cm² and can

be used continuously for up to 6 weeks as long as clinically indicated.

Intended Purpose Biatain Ibu Non-Adhesive Foam Dressing is intended for moist wound healing and exudate management of painful wounds; and is

indicated for a wide range of low to highly exuding wounds including acute wounds such as second degree burns, donor sites, postoperative wounds and traumatic wounds; and chronic wounds such as leg ulcers, pressure ulcers and non-infected diabetic foot ulcers. The dressing may reduce wound pain caused by tissue damage; and is suitable for use in combination with

compression therapy.

Non-adhesive dressings are suitable for use on fragile skin due to the absence of adhesive.

Variant information Size (cm) 10 x 10

Size (cm) 10 x 20 Size (cm) 15 x 15 Size (cm) 20 x 20

## **Specific Conditions**

No Specific Conditions included on Record

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