

# Askina® DresSil Active

## Product Data Sheet



### Administrative information

Legal Manufacturer	Winner Medical Co. Ltd. Winner Industrial Park, No 660 Bulong Road, Longhua District, 518109 Shenzhen, China
Importer	B. Braun Avitum AG, Schwarzenberger Weg 73-79, 34212 Melsungen, Germany
Product management	Wound Management
Last Update	30/05/2025

### Description, composition and properties of the device

Trade Name	Askina® DresSil Active
Reference	WIN5397510; WIN5391010; WIN5391110; WIN5391510; WIN5395210
Medical Class	Class IIb (MDR)
Description of the device	Askina® DresSil Active, a flex silicone foam dressing with border, 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.
Composition of the device	Polyurethane film backing Super absorbent fibers Non-woven fabric Polyurethane foam Silicone net Release film



## Key drivers & Indications

Key drivers	Good absorption and retention Maintains a moist wound environment Minimizes pain during dressing changes Easy application and removal Breathable Does not adhere to the wound bed
Intended Purpose	Askina® DresSil Active 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	Askina® DresSil Active 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 post operative blistering and may also be used as part of a prophylactic therapy to help prevent skin damage, e.g. pressure ulcers.
Precautions of use	<p>Askina® DresSil Active is contraindicated in:</p> <ul style="list-style-type: none"><li>• Check the wound for signs of infection before use, if infection occurs, see a health care professional.</li><li>• Do not use on the patients with a known hypersensitivity to the product itself or to its components.</li><li>• Do not use on third degree burns.</li></ul> <p>Warnings:</p> <ul style="list-style-type: none"><li>• Do not reuse. Reuse will cause cross-contamination.</li><li>• Do not use if package is damaged or open.</li><li>• Do not use together with oxidizing agents such as hypochlorite solutions or hydrogen peroxide.</li><li>• Due to the gentle adhesive used, do not use this dressing to secure other medical devices to the patient.</li><li>• If reddening or sensitization occurs discontinue use and see a healthcare professional.</li></ul>
IFU: Yes/No	Yes
Reusable/single use device	Single use device

## Sterilization process

Sterile: Yes/No	Yes
Sterilization method	Ethylene Oxide (EO)

## Conservation and storage conditions

Storage conditions	The product should be stored in dry conditions. Keep the product away from direct sunlight and keep dry.
Transport conditions	The product should be transported in dry conditions. Keep the product away from direct sunlight and keep dry.
Shelf life	3 years



## Safety in use

Technical: MRI, X-ray detectable

N/A

Biocompatibility

Biocompatibility studies showed the device has no indication of eliciting the following responses:

Cytotoxicity

Irritation or sensitization

Systemic Toxicity

Local effects after implantation

## Standards List

EN ISO 13485:2016

Medical devices – Quality management systems – Requirements for regulatory purposes

EN ISO 14971: 2019

Medical devices – Application of risk management to medical devices

EN ISO 20417:2021

Information supplied by the manufacturer of medical devices

EN ISO 15223-1: 2021

Medical devices – Symbols to be used with medical device labels, labelling and information to be supplied – Part 1: General requirements

ISO 15223-2: 2010

Medical devices – Symbols to be used with medical device labels, labelling, and information to be supplied – Part 2: Symbol development, selection and validation

EN ISO 10993-1: 2020

Biological evaluation of medical devices – Part 1: Evaluation and testing within a risk management process

EN ISO 10993-5: 2009

Biological evaluation of medical devices – Part 5: Tests for in vitro cytotoxicity

EN ISO 10993-10: 2013

Biological evaluation of medical devices – Part 10: Tests for irritation and skin sensitization

EN ISO 10993-11: 2018

Biological evaluation of medical devices – Part 11: Tests for systemic toxicity

EN ISO 10993-6: 2016

Biological evaluation of medical devices – Part 6: Tests for local effects after implantation

EN ISO 10993-11: 2018

Biological evaluation of medical devices – Part 11: Tests for systemic toxicity

EN ISO 11135: 2014/ A1: 2019

Sterilization of health-care products – Ethylene oxide – Requirements for the development, validation and routine control of a sterilization process for medical devices – Amendment 1: Revision of Annex E, Single batch release

ISO 10993-7: 2008/Amd 1: 2019

Biological evaluation of medical devices – Part 7: Ethylene oxide sterilization residuals – Amendment 1: Applicability of allowable limits for neonates and infants



Packaging & references			
Reference	Description	Size	Box Quantity
WIN5397510	Askina® DresSil Active	7,5 x 7,5 cm	10
WIN5391010	Askina® DresSil Active	10 x 10 cm	10
WIN5391110	Askina® DresSil Active	12,5 x 12,5 cm	10
WIN5391510	Askina® DresSil Active	15 x 15 cm	10
WIN5395210	Askina® DresSil Active	15 x 20 cm	10