

EXIT-PRO®

AG

S I L V E R

Description:

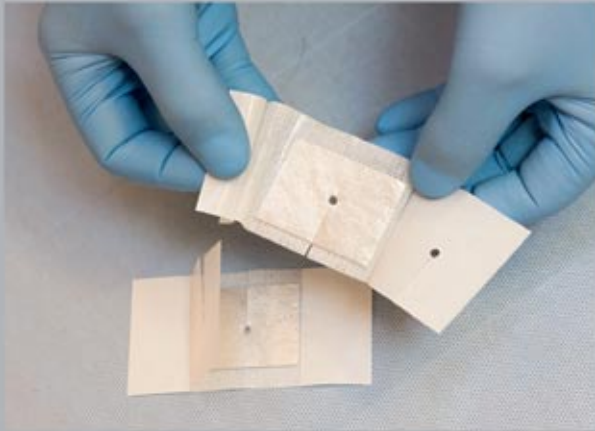
Non adherent silver dressing set on hypoallergenic plaster.

Indications:

Dressing of the intravascular devices connection sites to be used for the compression and the absorption of exudates.

Purpose:

The silver layer has an immediate and long term antibacterial action helping to reduce the risk of local infection of the exit-site, of the tunnel or systemic (catheter-correlated sepsis) related to the use of intravascular devices. It helps to prevent the irritation caused by the mechanical rubbing of the catheter over the skin. Besides, compared to other antiseptics, the silver ions of *EXIT-PRO-Ag* in a $<0,02\mu\text{g}/\text{cm}^2$ concentration, are well tolerated.



Pat Pending

REFERENCES Data and technical specification can be changed without notice

NON ADHERENT ADHESIVE SILVER DRESSING

CODE	PAD Measure cm	PLASTER Measure cm	Quantity blister pack	Package box
S0810/AG	10x5	3,5x4,5	1	250
S0815/AG	15x5			
S0808/AG	8x2,5	2,5x3,5	2	250

Sperimental evidences

The antibacterial action *EXIT-PRO-Ag* pads has been tested.

The results are shown in the down below diagram.

Antibacterial action of

Biochem test report n° 2498/08

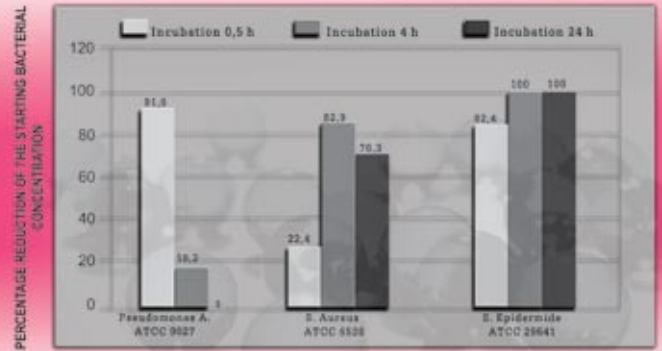
Percentage reduction of the starting bacterial concentration

incubation 0,5h- incubation 4h- incubation 24h

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Method of the analysis

A standardised bacterial culture (minimum starting concentration of 1×10^6 cfu/ml) is placed onto the sample (pad with silver) and control (pad without silver). The count of the bacterial colonies grown is made immediately after sowing (0,5 h), after 24 h and 48 h incubation period. The results are expressed as percent bacterial reduction in *EXIT-PRO-Ag* sample vs. control.



Direction for use

- 1 Put the dressing over the catheter exit-site and hold in place by pressing on the plaster
- 2 Change dressing at every session

Classification: Class III medical device
Biocompatibility: The material in contact with the skin has passed the biocompatibility tests according to UNI EN ISO 10993 standards
Sterilisation: γ - ray
Expiry: 5 years