

Medicomp® Drain sterile

General Product Description

Sterile non-woven slit swabs made of a mixture of viscose and polyester fibres, the part of viscose is between 67 and 70% the remaining part is made of polyester fibres available in different sizes with one cut in Y form. The sterile swabs are packed in paper-film-pouches.

Indication

Slit non-woven swabs can be used to provide padding and protection, when a needle or similar device penetrating the skin (e.g. cannula, catheter) is inserted into the human body to access veins for administration of a medicinal product or insertion of a medical device into the human body for absorption of exuding body fluids.

Reference Numbers

REF	name	size [cm]	grammage	number of ply	pieces per pouch	pouches per box
4215338	Medicomp® Drain sterile	7.5x7.5	30	6	2	25
4215356	Medicomp® Drain sterile	10x10	30	6	2	25



Residual Risks, Contra-Indications and Undesirable Side Effects, Warnings

Not known

Sterile Device

The sterile packaging is in accordance with DIN EN ISO 11607.

Sterilization with steam according to DIN EN ISO 17665-1 with validated process

Instruction	Symbol acc. to EN ISO 15223-1
Do not use if package is damaged	
Do not resterilize	

Single Use Device

Reusing a single-use medical device is dangerous. Reprocessing devices, in order to reuse them, may seriously damage their integrity and their performance. Information available on request.

Product Disposal

To minimize the risk of potential infection hazards, or environmental pollution, disposable components of Medicomp® Drain sterile should follow disposal procedures according to applicable and local laws, rules, regulations and infection prevention standards.

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Incident Reporting

For a patient/user/third party in the European Union and in countries with identical regulatory regime (Regulation 2017/745/EU on Medical Devices); if, during the use of this device or as a result of its use, a serious incident has occurred, please report it to the manufacturer and/or its authorised representative and to your national authority.

Product Performance Characteristics

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property	value
Area weight	approx. 30 g/m ²
Water absorption capacity	approx. 7 g/g
Optical brighteners	none
Chemical binders	none

Material Characteristics

Non-woven

Mixture of viscose and polyester

Labelling

Lot-No. with 9-digit code

e.g.:



9

999

01

00

1

explanation [9 = 2019]

definition year

[999-000]

serial production order

for internal purposes only

Manufacturing date

e.g.:



2019
year

04
month

01
day

Expiry date

e.g.:



2024
year

04
month

01
day

Shelf Life: 5 years

Medical Device



Unique Device Identification (UDI)



Single Sterile Barrier System



Single sterile barrier system

Latest Date of Revision: 2020-08-04