

See-through packaging made of paper and film VP code/quality: MM70-1/PP50-1

1. Product description

Sterile barrier system for medical disposables and reprocessed medical devices.

2. Sterilisation suitability

Steam, ethylene oxide and formaldehyde sterilisation

- Indicator change

STEAM:	pink to brown
EO:	light blue to yellow
FORM:	purple to green



3. Normative requirements

- The paper complies with DIN EN ISO 11607 Part 1 and DIN EN 868 Part 3.
- The film complies with DIN EN ISO 11607 Part 1 and DIN EN 868 Part 5 Section 4.2.2.1 to 4.2.2.5.
- The packaging material complies with DIN EN ISO 11607 Part 1 and Part 2.
- The biocompatibility and toxicological properties were tested and evaluated for cytotoxicity according to ISO 10993 Part 5.
- Microbial barrier for moisture and when dry according to DIN 58953 Part 6.
- The process indicators comply with DIN EN ISO 11140 Part 1.

Any standards is applicate in its current version.

4. Statutory requirements

Our see-through pouches and see-through reels are accessories to medical device and as such medical devices of class I "non-sterile". This classification is according to annex VIII, rule 1 of the Medical Device Regulation (EU) 2017/745.

As an indication of compliance with the regulation the label of the shipping carton carries the CE marking.

In addition, our products comply with the following regulations and directives:

- Regulation (EU) No. 10/2011
- Regulation (EC) 1935/2004
- Regulation (EC) No. 1907/2006
- Directive 2011/65/EU Art. 4 and Annex II
- Directive 94/62/EC

The sums of the concentrations of lead, cadmium, mercury and hexavalent chromium (Cr VI) are below 100 ppm.

5. Substances

Our products do not contain natural latex, PVC, Bisphenol A, Colophony, TSE/BSE risk materials, SVHC/CP65-Substances and Phthalates as part of the formulations.

6. Product properties

Paper

The white medical paper is physiological harmless, free of foreign substances, and referred to as **elemental chlorine-free (ECF)**. The mass fraction of the paper in chloride, expressed as sodium chloride is far below the required standard DIN EN 868 Part 3 Point 4.2.5.

The specially treated surface enables a firm seal against the film.

Inspection characteristic	Typical value	Unit	Standards
Grammage	70	g/m ²	DIN EN ISO 536
Tensile strength MD dry	120	N/15 mm	DIN EN ISO 1924-2
Tensile strength CD dry	60	N/15 mm	DIN EN ISO 1924-2
Tensile strength MD wet	20	N/15 mm	DIN ISO 3781
Tensile strength CD wet	10	N/15 mm	DIN ISO 3781
Burst strength dry	350	kPa	DIN EN ISO 2758
Burst strength wet	70	kPa	DIN ISO 3689
Tear strength MD	650	mN	DIN EN 1974
Tear strength CD	750	mN	DIN EN 1974
Air porosity Bendtsen	680	ml/min.	ISO 5636-3
Water resistance	25	sec.	DIN EN 868-3
pH value	6.5		ISO 6588-2
Chlorine content	< 0.05	%	DIN EN 868-3
Sulphate content	< 0.25	%	DIN EN 868-3
Pore size	20	µm	DIN EN 868-3

Film

The 12/40 PET/PP film is blue, physiological harmless and heat-sealable against medical paper.

Inspection characteristic	Typical value	Unit	Standards
Configuration	12/40 PET/PP		
Thickness	54	µm	DIN 53370
Grammage	54	g/m ²	Internal
Tear strength CD	> 35	N/15mm	ASTM-D 882
Tear strength MD	> 30	N/15mm	ASTM-D 882
Elongation at breakage CD	> 70	%	ASTM-D 882
Elongation at breakage MD	> 80	%	ASTM-D 882

7. Seal seam design

The seal seam allows the sterile barrier system to be easily opened. The seal seam consists of three parallel seal lines (grooved seal seam).

The minimum width of the seal(s) is 6 mm. In grooved seals the sum of the rib width of ribs is minimum 6 mm.

The minimum value of the seal strength in the sterilisation process with STEAM is 1.5 N per 15 mm, and with other sterilisation processes used in healthcare 1.2 N per 15 mm.

8. Peel direction

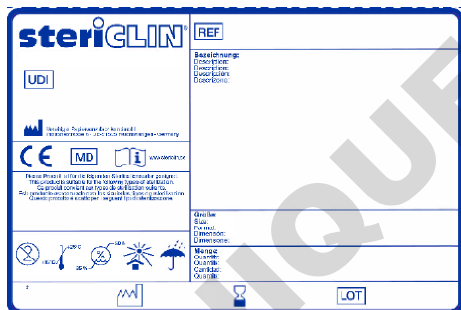


The correct peel direction is printed on the see-through pouch or see-through reel to ensure its safe, fibre-free opening.

9. Identification

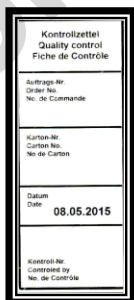
See-through pouch

Carton label

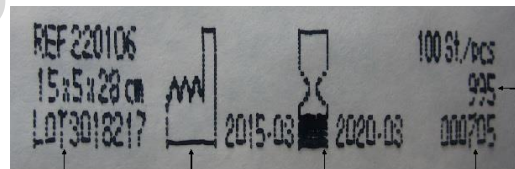


Box No. Production Expiry date LOT-No.

Control document for pouches without secondary packaging



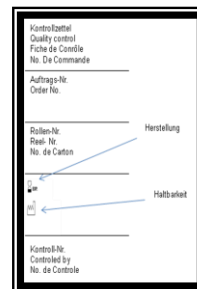
Printed on the secondary packaging



LOT-No. Production Expiry date Package No.

Personnel No.

Control document for see-through reels



10. Disposal

The sterile barrier systems are composites or mixed packaging (waste codes 15 01 05 and 15 01 06) and must be disposed of in the non-recyclable waste.

11. Assortment

See-through pouch without gusset

<https://www.stericlin.de>

See-through reel without gusset

<https://www.stericlin.de>

See-through pouch with gusset

<https://www.stericlin.de>

See-through reel with gusset

<https://www.stericlin.de>

12. Field of application

This Technical Data Sheet refers to the above-mentioned product group/quality and applies until the next revised edition. Other product-related documents are available on request if required.

13. Additional information

The information contained in this Technical Data Sheet (TDS) is based on our current knowledge and experience.

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