



Processing instructions of NOBAMED Paul Danz AG  
for NOBA® medical products made of **non-woven**  
that are to be sterilized before use.

**The product must be sterilized before use. We confirm the one-time sterilizability with a procedure mentioned under the section 'sterilization'.**

<b>Instructions</b>	
<b>Initial treatment at the place of use (storage and transport)</b>	The storage and transport conditions and the shelf-life of the non-sterile product are indicated on the package
<b>Preparation of sterilization</b>	Remove and discard the packaging of the product
<b>Cleaning</b>	Not applicable
<b>Disinfection</b>	Not applicable
<b>Drying</b>	Not applicable
<b>Control and verification</b>	The validation of the sterilization process and the product fall under the responsibility of the processor
<b>Packaging</b>	Suitable validated sterile packaging
<b>Sterilization</b>	For example <ul style="list-style-type: none"><li>• with moist heat according to DIN EN ISO 17665 or as per requirements of the European Pharmacopoeia with 15' to 20', 121°C to 134°C, 2 bar</li><li>• or with one-time sterilizability with an ethylene oxide sterilization method in compliance with ISO 11135-1, e. g. max. 60 °C and max. 90 % RH. A sterilization below these limit values does not affect the products' functionality, nor do pressure or the rate of pressure change.</li></ul>
<b>Storage and transportation</b>	For the product the storage and transport conditions indicated on the original packaging are applicable.
<b>Additional information</b>	NOBA® medical products made of non-woven are disposable products and cannot be reprocessed for product-related reasons. This does not refer to the process of first-time sterilization, if required before use. Our products are produced under controlled conditions. The bioburden of non-sterile products is $<10^2$ cfu/g.

The above mentioned instructions have been compiled in accordance with EN ISO 17664-1 and have been validated as SUITABLE for the processing of NOBA® medical products made of non-woven. It falls under the processor's responsibility to make sure that the actual process (with the equipment, materials and staff in the processing department) is adequate to achieve the desired results. This necessitates a validation and routine monitoring of the process on site.

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