

» Declaration of clearance «

Under the MDR (Medical Device Regulation, EU Regulation 2017/745), materials used for the production of medical products must be provided, amongst others, with a declaration of clearance with respect to their toxicological properties.



nora° products are free of constituents posing toxicological and carcinogenic risks as set down under EU Regulation 2017/745.

The closed-cellular surface structure of **nora**® materials also minimises the risk of harm to health in the form of germ and bacterial colonies and facilitates complete hygienic cleaning and disinfection.

Owing to their properties, **nora*** materials are suitable for the processing of Class 1 medical products.

nora* products undergo regular testing at renowned testing institutes and are certified to bear the following quality labels:

- dermatological seal of the Dermatest institute
- SG tested for harmful substances of the testing and research institute PFI Pirmasens

These requirements are fulfilled as a result of continuous production monitoring by an extensive quality management system. The ISO 9001:2015 certification safeguards unvarying quality in the manufacture, sales, and marketing of **nora*** products. This is supplemented with an **environmental management system** complying with ISO 14001:2015.

Confirmed in January 2025. Valid from 01/01/2025 to 31/12/2026.

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