

## STATEMENT ON COMPLIANCE TO REGULATIONS ON MEDICAL USE

We confirm that this product fulfils the requirements on materials used for the manufacturing of articles or components of articles intended for medical use as described in:

**Council of Europe** Material complies with the following European Pharmacopoeia monographs:

Monograph 3.1.3. Polyolefins

Monograph 3.1.6. Polypropylene for containers and closures for parenteral

preparations and ophthalmic preparations

Tests are made according to the current Pharmacopoeia edition at the time

of the testing: 10th edition (2020), and supplement 10.2 (07/2020).

Monograph 3.2.2. Plastic containers and closures for pharmaceutical use: This monograph relates specifically to the <u>container and closure system</u> and does not contain tests that are required for compliance assessment of the raw materials used for said container and closures. The composition of the

product is in compliance with this monograph.

**Germany** The product follows the VDI 2017 Guideline on "Medical Grade Plastic" that

covers the requirements for change management, quality management,

supply security and support for regulatory requirements.

**USA**Material has passed the following United States Pharmacopeia tests:

Biological reactivity tests - in vitro <87>: Cytotoxicity (Elution test)

Biological reactivity tests - in vivo <88>: Class VI - 70 °C

Plastic materials of construction <661.1>: Identification, physicochemical and extractable metals (as listed in the chapter) tests. Plastic additive tests are

done according to Borealis' internal methods.

Tests are made according to the current Pharmacopeia edition at the time of

the testing (USP 39).

**Additional testing** Material has been tested according to the following ISO 10993 biological

tests, in the extent applicable for polymer pellets:

Cytotoxicity

Acute systemic toxicity

Skin irritation (intracutaneous reactivity)

Dermal sensitization Hemocompatibility

Tests are made according to the current ISO 10993 edition at the time of the

testing (2016).

**Elemental impurities** During the manufacturing process of this product, we neither use nor

intentionally incorporate any Class 1, 2A, 2B or 3 elements as listed in the

ICH Q3D(R1) Guideline on Elemental Impurities (March 2019).

Bormed is a trademark of the Borealis group.





**DMF number** Material has been assigned the FDA Drug Master File number(s):

DMF 017929

Additional information If a customer wishes to take advantage of the pre-notice period in case of

deletion or modification of Bormed grades, such pre-notice period needs to be

included in Technical Delivery Specifications.

This edition of the document supersedes any previous editions.

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Prepared by Borealis, Group Product Stewardship / Sofia Montalvão

## **Disclaimer**

The product(s) mentioned herein are not intended for use as medical implant material or implantable medical devices and we do not support their use for such applications.

To the best of our knowledge, the information contained herein is accurate and reliable as of the date of publication; however we do not assume any liability whatsoever for the accuracy and completeness of such information.

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