

EU Quality Management System Certificate

Certificate no.
3558GB448240808

Final Assessment Report no.
3558AU22F

Effective date
2024-08-08

Expiry date
2026-07-28

This is to certify that the quality system of
HÄLSA Pharma GmbH
Maria-Goeppert-Straße 5, 23562 Lübeck, Germany
SRN: DE-MF-000007407

For design, production, and final product inspection/testing of
Medical devices/groups of medical devices listed on the following pages

Has been assessed and found to comply with respect to
**The conformity assessment procedure described in Annex IX,
Chapters I and III of Regulation (EU) 2017/745 on Medical Devices**

Any applicable limitations for certain medical devices are included in the following list or recorded in the final assessment report. This certification is subject to surveillance by DNV MEDCERT.

Place and date
Hamburg, 2024-08-08

For the issuing office
DNV MEDCERT GmbH – Notified Body 0482
Pilatuspool 2, 20355 Hamburg, Germany



Benannt durch/Designated by
Zentralstelle der Länder
für Gesundheitsschutz
bei Arzneimitteln und
Medizinprodukten
www.zlg.de
BS-MDR-096

The certificate is only valid when provided entirely with all of its pages. To verify the validity of this certificate, contact Medcert-Info@dnv.com


Lorenz Runge
Director Certification Body



Certificate no.: 3558GB448240808
Place and date: Hamburg, 2024-08-08

Preceding certificate

Certificate no.	Issue date	Identification of changes
3558GB448220714	2022-07-14	WO-009807, Addition of MDN 1213
3558GB448240227	2024-02-27	WO-009830, WO-010350

Sites covered by this certificate

HÄLSA Pharma GmbH, Maria-Goeppert-Straße 5, 23562 Lübeck, Germany
HÄLSA Pharma GmbH, Am Mittelhafen 56, 48155 Münster, Germany



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Products covered by this certificate

Class IIa medical devices

Category	EMDN code	Medical devices/groups of medical devices
MDN 1213	Q019003	Dental floss and other devices for oral hygiene (for professional use)

Class III medical devices

For placing on the market of class III medical devices covered by this certificate, an additional EU Technical Documentation Assessment Certificate according to Annex IX Chapter II of Regulation (EU) 2017/745 is required, which also contains the exact determination of medical devices covered by certification.

Category	Medical devices/groups of medical devices
MDN 1204	Non-active non-implantable devices for wound and skin care
MDN 1213	Non-active non-implantable devices composed of substances to be introduced into the human body via a body orifice or the dermal route