





Product Service

EU Technical Documentation Assessment Certificate (MDR)

Pursuant to Regulation (EU) 2017/745 on Medical Devices, Annex IX Chapter II (Implantable Class IIb Devices and Class III Devices)

No. G70 027023 0171 Rev. 01

Manufacturer: **CROMA-PHARMA GmbH**

> Industriezeile 6 2100 Leobendorf **AUSTRIA**

SRN Manufacturer - AT-MF-000013284

The Certification Body of TÜV SÜD Product Service GmbH certifies that the manufacturer has drawn up and presented a Technical Documentation according to Annex II and III of the Regulation (EU) 2017/745 on medical devices. Details on devices covered by the Technical Documentation are described on the following page(s).

The Report referenced below summarises the result of the assessment and includes reference to relevant CS, harmonized standards and test reports. The technical documentation assessment included an assessment of the clinical evaluation assessment.

The conformity assessment has been carried out according to Annex IX chapter II of this regulation with a positive result. Changes to the approved device, where such changes could affect the safety and performance of the device or the conditions prescribed for use of the device, shall require approval from the notified body TÜV SÜD Product Service GmbH.

In order to place the devices on the market with CE-marking, an EU Quality Management System Certificate pursuant to Annex IX chapters I and III is necessary in addition to this EU Technical Documentation Assessment Certificate. All applicable requirements of the Testing, Certification, Validation and Verification Regulations TÜV SÜD Group have to be complied with. For details and certificate validity see: www.tuvsud.com/ps-cert?q=cert:G70 027023 0171 Rev. 01

Report No.: 713342326

Preceding Certificate No.: G70 027023 0171 Rev. 00

Valid from: 2024-09-09 Valid until: 2029-08-21

Date of Initial Issuance: 2023-12-20

Christoph Dicks

Head of Certification/Notified

Body



Product Service

EU Technical Documentation Assessment Certificate (MDR)

Pursuant to Regulation (EU) 2017/745 on Medical Devices, Annex IX Chapter II (Implantable Class IIb Devices and Class III Devices)

No. G70 027023 0171 Rev. 01

Classification: Class III (Annex XVI)

Device Group: P900402 - RESORBABLE FILLING AND RECONSTRUCTION

DEVICES

Basic UDI-DI: 9003502BIMDVOLIDO001SX

Intended Purpose: The intended aesthetic purpose of the viscoelastic implant is to

add volume.

Device with non-medical intended purpose according to article 1

(2)

Device(s): saypha volume Lidocaine

Apriline Forte Lidocaine Beaubella Strong Lido

Definisse restore filler + Lidocaine DERMALSTYLE forte Lidocaine

Duraybo Strong Lido Faysia Forte Lido JALOR Sweet Kiss Pluryal Volume Lidocaine Princess VOLUME Lidocaine

Quelvez Forte Lido RENNOVA LIFT LIDO XCELENS extra 4

ZFill deep+



Product Service

EU Technical Documentation Assessment Certificate (MDR)

Pursuant to Regulation (EU) 2017/745 on Medical Devices, Annex IX Chapter II (Implantable Class IIb Devices and Class III Devices)

No. G70 027023 0171 Rev. 01

The validity of this certificate ./. depends on conditions and/or is limited to the following:

Revision History:

Rev.	Dated	Report	D
00	2023-12-20	713265756	Ir
01	2024-09-09	713342326	Α

Description nitial issuance Amended: Other

Spelling change of saypha