

## **EC** Certificate

## Directive 93/42/EEC Annex II, excluding Section 4 Full Quality Assurance System **Medical Devices**

Registration No.: HD 60148183 0001

Report No.:

21270380 007

Manufacturer:

DIRINCO B.V. Ketelmeer 1 5347 JX Oss **Netherlands** 

**Products:** 

Anticoagulant for use as catheter lock

(see attachment for products included)

Replaces Certificate, Registration No.: HD 60124779 0001

**Expiry Date:** 

2024-05-26

The Notified Body hereby declares that the requirements of Annex II, excluding section 4 of the directive 93/42/EEC have been met for the listed products. The above named manufacturer has established and applies a quality assurance system, which is subject to periodic surveillance, defined by Annex II, section 5 of the aforementioned directive. For placing on the market of class III devices covered by this certificate an EC design-examination certificate according to Annex II, section 4 is required.

Effective Date:

2020-03-26

Date:

2020-03-26

**Notified Body** 

Roland Gruber

TÜVRheinle

TÜV Rheinland LGA Products GmbH - Tillystraße 2 - 90431 Nü TÜV Rheinland LGA Products GmbH is a Notified Body according to Directive 93/42/PEC concerning medical devices with the identification number 0197.



Doc. 1/1, Rev. 0

## TÜV Rheinland LGA Products GmbH Tillystraße 2, 90431 Nürnberg

Attachment to Certificate

Registration No.:

HD 60148183 0001

Report No.:

21270380 007

Manufacturer:

DIRINCO B.V. Ketelmeer 1 5347 JX Oss Netherlands

## Products included:

Anticoagulant, antimicrobial and reducing biofilm formation for use as catheter lock

- CitraLock 30% and 46,7%
- 5 ml in Polyethylene vials

Anticoagulant for use as catheter lock CitraLock 4%

- 5 ml in Polyethylene vials

Date: 2020-03-26

Roland Grubers Roland