

## **EC** Certificate

**Certificate Number:** 

**DGM - 673** 

This is to certify that the quality system of:

## Vitrolife A/S

(Unisense Fertilitech A/S, Fertilitech A/S) Jens Juuls Vej 20 8260 Viby J Denmark

has been approved in conformity with the requirements of

## Annex V, section 3.2 - Production quality assurance

of Council Directive 93/42/EEC concerning medical devices as transposed into Danish law.

The certificate covers the following activities:

Manufacture and final inspection of IVF incubators and related accessories in class lla

The EC certificate is valid provided that the quality system continues to conform to the above-mentioned scope and provided that the company does not introduce substantial changes to the quality system without the approval of Presafe Denmark A/S. This EC certificate is issued pursuant to the Presafe Denmark A/S terms and conditions for the certification of medical devices and entitles the manufacturer to affix the CE mark. The certificate is based on successful audit of the manufacturer. The manufacturer is subject to periodical audits as per the Directive.

Heidi Jørgensen Authorized person

For Presafe Denmark A/S

Date of issue:

2015-04-24

Expires:

2017-06-29

Initial date of issue: 2009-06-29

Reference:

aur5a1503v190f712

## Presafe Denmark A/S

Notified Body, Identification No. 0543 Tuborg Parkvej 8, 2900 Hellerup, Denmark





The following product families in class IIa are covered by the certificate:

EmbryoScope<sup>™</sup> time-lapse system

consisting of:

EmbryoScope<sup>™</sup> time-lapse incubator

EmbryoSlide® culture dish

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Certificate type: EC Certificate



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