



Product Service

EC Certificate

EC Design-Examination Certificate

Directive 93/42/EEC on Medical Devices (MDD), Annex II (4)
(Devices in Class III)

No. G7 16 09 74018 015

Manufacturer: CERUS Corporation

2550 Stanwell Drive
Concord CA 94520
USA



EC-Representative: Cerus Europe B.V.

Stationsstraat 79-D
3811 MH Amersfoort
THE NETHERLANDS

Product: Blood Processing Devices
Pathogen Inactivation Disposables

The Certification Body of TÜV SÜD Product Service GmbH declares that a design examination has been carried out on the respective devices in accordance with MDD Annex II (4). The design of the devices conforms to the requirements of this Directive. For marketing of these devices an additional Annex II certificate is mandatory. See also notes overleaf.

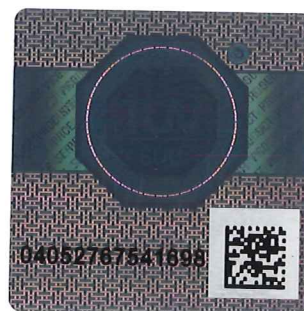
Report no.: 713082447

Valid from: 2016-09-29

Valid until: 2021-09-14

Date, 2016-09-29

Stefan Preiß



TÜV SÜD Product Service GmbH is Notified Body with identification no. 0123

Page 1 of 2



Product Service

EC Certificate

EC Design-Examination Certificate

Directive 93/42/EEC on Medical Devices (MDD), Annex II (4)
(Devices in Class III)

No. G7 16 09 74018 015

Model(s): INTERCEPT Blood System for Plasma
(Amotosalen Photochemical Treatment)

Parameters: INTERCEPT Processing Set for Plasma, INT31

Facility(ies): CERUS Corporation
2550 Stanwell Drive, Concord CA 94520, USA

Cerus Corporation
2411 Stanwell Drive, Concord CA 94520, USA

Cerus Europe B.V.
Stationsstraat 79-D, 3811 MH Amersfoort, THE NETHERLANDS