



Directive 93/42/EEC on Medical Devices, Annex V

No. CE 00362

Issued To: Becton, Dickinson and Company

Belliver Industrial Estate

Belliver Way

Roborough, Plymouth

PL6 7BP

United Kingdom

In respect of:

The manufacture of sterile safety lancets, blood collection needles and syringes with preattached needles, to be used for the collection of blood for in vitro diagnostic examination.

on the basis of our examination of the quality assurance system under the requirements of Council Directive 93/42/EEC, Annex V. The quality assurance system meets the requirements of the directive. For the placing on the market of class IIb and class III products an Annex III certificate is required.

For and on behalf of BSI, a Notified Body for the above Directive (Notified Body Number 2797):

Gary E Slack, Senior Vice President Medical Devices

Gay C Stade

First Issued: **1994-12-22** Date: **2019-11-28** Expiry Date: **2024-05-26**

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Validity of this certificate is conditional on the quality system being maintained to the requirements of the Directive as demonstrated through the required surveillance activities of the Notified Body. This approval excludes all products designed and/or manufactured by a third party on behalf of the company named on this certificate, unless specifically agreed with BSI.

This certificate was issued electronically and is bound by the conditions of the contract.





Supplementary Information to CE 00362

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Product Information Table

| Number | Device Name | Intended Purpose per IFU | | |
|-----------|--|---|--|--|
| Class IIa | | | | |
| MD 0106 | BD Sentry Safety Lancet | Sterile, single-use, micro-collection lancet used to perform finger-stick punctures in adults and children for sampling capillary blood. | | |
| MD 0106 | BD Preset & Preset Eclipse Arterial Blood Critical Care Collection Syringes (with pre-attached needle) | Sterile, single use device to be used for the collection, primary containment and preservation of blood specimens derived from the human body for the purposes of IVD examination | | |
| MD 0106 | BD Vacutainer PrecisionGlide Multiple Sample Blood Collection Needle | Sterile single use medical device intended to be used for the sampling of venous blood derived from the human body for the purposes of for IVD examination | | |
| MD 0106 | BD Vacutainer Eclipse Signal Blood Collection Needle | Sterile, single use device for the collection of multiple blood samples into evacuated blood collection tubes for the purpose of in vitro testing | | |

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| Number | Device Name | Intended Purpose per IFU | | |
|---------|---|---|--|--|
| Class I | | | | |
| MD 0106 | BD Vacutainer Blood Collection Needle Holder | One Use Holder intended to be used in conjunction with the BD Vacutainer® range of blood collection needles, blood collection sets and luer adapter in order to facilitate the insertion of the needle into the patient's vein and to help guide the evacuated blood collection tube onto the non-patient (NP) end of the needle during the blood collection process. | | |

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Directive 93/42/EEC on Medical Devices, Annex V

List of Significant Subcontractors

Recognised as being involved in services relating to the product covered by:

Certificate No: **CE 00362** Date: 2019-11-28

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Subcontractor:

Becton Dickinson Switzerland Sarl

Terre Bonne Park - A4 Route de Crassier - 17

1262 Eysins Switzerland

HTL-Strefa S.A.

Ul. Adamówek 7

95-035 Ozorków

Poland

HTL-Strefa S.A. (Leczyca)

UI. Lotnicza 21h

Poland

99-100 Leczyca

Sterigenics Belgium (Fleurus) SA Zoning Industriel de Fleurus

Avenue De L'Esperance Fleurus, Hainut

B-6220

Belgium

Service(s) supplied

EU Representative

Finished Device Supplier

Manufacture

Gamma Sterilization

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Subcontractor:

Service(s) supplied

Synergy Health Radeberg GmbH Juri-Gagarin Str. 15 D-01454 Radeberg Germany **Gamma Sterilization**

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EC Certificate - Production Quality Assurance Certificate History

Certificate No:

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Date:

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| Date | Reference Number | Action |
|------------------|---------------------|---|
| 22 December 1994 | | First issue. |
| 12 August 1997 | | Change of post code on address. |
| 10 December 1997 | | Change of Notified Body (BSI) address. |
| 22 February 2001 | | "with and without anticoagulant for diagnostic purposes" added to the scope, also removal of Griffith Microsciences (Somercotes) from sub-contractor list. |
| 11 March 2004 | | Company name change to Becton Dickinson & Company, 5 year renewal, and addition of IBA S & I Limited, IBA Medisus S/A, Hepartex SA and Celsus Labs. Inc. |
| 30 November 2009 | | Certificate renewal. Removal of 'Hepartex S.A', 'Celsus Laboratories, Inc.' and 'IBA S & I Limited' as subcontractors. Change of subcontractor name 'IBA Mediris S.A' to 'Sterigenics Belgium (Fleurus) SA and rewording of scope. |
| 18 December 2014 | 8215528 | Change of scope wording from 'syringe and needles' to 'sterile collection needles and syringes with pre-attached needles' Certificate renewal. |

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|---------------|---------------------|--|
| 10 March 2017 | 8693238 | Scope extension for sterile safety lancets (traceable to CE 583593). |
| | | Addition of subcontractors HTL-Strefa, Poland as finished device supplier and Synergy Health Radeberg for gamma sterilisation of safety lancets. |
| 05 March 2019 | 7779292 | Traceable to NB 0086. |
| Current | 9769232 | Certificate Renewal. |
| | | Addition of product information table. |
| | | Addition of subcontractor BD Switzerland Sarl as EU Authorised Representative & HTL-Strefa S.A. Leczyca) for manufacture. |

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