

## EC DECLARATION OF CONFORMITY

<b>Manufacturer:</b>	<b>Becton, Dickinson and Company</b> Belliver Industrial Estate, Belliver Way, Roborough, Plymouth, PL6 7BP, United Kingdom	
<b>Manufacturing Site(s):</b>	<b>HTL-Strefa S.A.</b> Ul. Adamówek 7, 95-035 Ozorków Poland	
<b>Products:</b>	<b>Catalogue number</b> 369523 369528	<b>Device name</b> BD Sentry™ Safety Lancet BD Sentry™ Safety Lancet
<b>Classification:</b>	Class IIa	
<b>Conformity Assessment Route:</b>	Conformity is established through application of the procedures described in Annex V and Annex VII of the European Medical Devices Directive 93/42/EEC.	
<b>GMDN:</b>	61578 – Manual blood lancing device, single-use	
<b>Notified Body:</b>	British Standards Institution (BSI), Kitemark Court, Davy Avenue, Knowlhill, Milton Keynes, MK5 8PP, UK. (NB Number 0086)	
<b>CE Certificate Number:</b>	00362	
<b>Date of issue of original CE Certificate:</b>	08 March 2012 (Original CE Certificate CE 583593)	

We herewith declare that the above mentioned products meet the provisions of the Council Directive 93/42/EEC of 14 June 1993 concerning medical devices. All supporting documentation is retained under the premises of the manufacturer.

<b>List of Harmonised Standards:</b>
<p><b>EN ISO 13485:2012</b> Medical devices - Quality management systems - Requirements for regulatory purposes, <b>EN ISO 14971:2012</b> Medical devices - Application of risk management to medical devices, <b>EN ISO 11737-1:2006</b> Sterilization of medical devices. Microbiological methods. Determination of a population of microorganisms on products, <b>EN ISO 11737-2:2009</b> Sterilization of medical devices. Microbiological methods. Tests of sterility performed in the definition, validation and maintenance of a sterilization process, <b>EN ISO 11137-1:2015</b> Sterilization of health care products - Radiation - Part 1: Requirements for development, validation and routine control of a sterilization process for medical devices, <b>EN ISO 11137-2:2015</b> Sterilization of health care products. Radiation. Establishing the sterilization dose, <b>EN ISO 15223-1:2016</b> Medical devices - Symbols to be used with medical device labels, labelling and information to be supplied - Part 1: General requirements, <b>EN 1041:2008 +A1:2013</b> Information supplied by the manufacturer of medical devices, <b>EN ISO 11607-1:2009+A1:2014</b> Packaging for terminally sterilized medical devices. Requirements for materials, sterile barrier systems and packaging systems, <b>EN ISO 10993-1:2009</b> Biological evaluation of medical devices. Evaluation and testing within a risk management process, <b>EN ISO 10993-2:2006</b> Biological evaluation of medical devices. Animal welfare requirements, <b>EN ISO 10993-3:2014</b> Biological evaluation of medical devices - Part 3: Tests for genotoxicity, carcinogenicity and reproductive toxicity, <b>EN ISO 10993-4:2009</b> Biological evaluation of medical devices - Part 4: Selection of tests for interactions with blood, <b>EN ISO 10993-5:2009</b> Biological evaluation of medical devices. Tests for in vitro cytotoxicity, <b>EN ISO 10993-11:2009</b> Biological evaluation of medical devices. Tests for systemic toxicity, <b>EN ISO 10993-12:2012</b> Biological evaluation of medical devices - Part 12: Sample preparation and reference materials, <b>EN ISO 10993-13:2010</b> Biological evaluation of medical devices - Part 13: Identification and quantification of degradation products from polymeric medical devices <b>EN ISO 10993-15:2009</b> Biological evaluation</p>



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of medical devices - Part 15: Identification and quantification of degradation products from metals and alloys **EN ISO 10993-17:2009** Biological evaluation of medical devices - Part 17: Establishment of allowable limits for leachable substances **EN ISO 10993-18:2009** Biological evaluation of medical devices - Part 18: Chemical characterization of materials **EN 556-1:2001** Sterilisation of medical devices - Requirements for medical devices to be designated "STERILE" - Part 1: Requirements for terminally sterilised medical devices

**List of Non-Harmonised Standards:**

**ISO 14644-1:2015** Cleanrooms and associated controlled environments - Part 1: Classification of air cleanliness, **ISO 10993-4:2017** Biological evaluation of medical devices - Part 4: Selection of tests for interactions with blood, **ISO 10993-6:2016** Biological evaluation of medical devices - Part 6: Tests for local effects after implantation, **EN ISO 10993-10:2013** Biological evaluation of medical devices - Part 10: Tests for irritation and skin sensitization, **EN ISO 12048:2002** Packaging - Complete, filled transport packages - Compression and stacking tests using a compression tester, **EN 22248:2001** Packaging - Complete, Filled Transport Packages - Vertical Impact Test By Dropping, **EN 24180-2:2002** Guide to compilation of performance test schedules for complete, filled transport packages. Quantitative data, **EN 28768:2002** Packaging - Complete, Filled Transport Packages - Toppling Test, **EN ISO 2234:2007** Packaging -- Complete, filled transport packages and unit loads -- Stacking tests using a static load, **EN 60068-2-31:2010** Environmental testing. Tests. Test Ec. Rough handling shocks, primarily for equipment-type specimens

SIGNED FOR AND ON BEHALF OF: Becton, Dickinson and Company

PLACE, DATE OF ISSUE: Plymouth, 01<sup>st</sup> February 2018

Signature: \_\_\_\_\_

Lorna Darrock

EMEA Regulatory Affairs and Compliance Manager

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### **VERSION HISTORY**

Current Version Prepared By: Matthew Piska

<b>REV.</b>	<b>Version Description</b>
A	Transferred from QDMS to ECC – Version number remained 1
B	Transfer into new Medical Declaration of Conformity Template (MED-RA-001C). Replaced CE Certificate 583593 with CE Certificate 00362. Replaced obsolete GMDN code 37466 with 61578
C	Update to harmonised and non-harmonised standards list