Belliver Industrial Estate Belliver Way, Roborough Plymouth, PL6 7BP United Kingdom tel: +44(0)1752 701281 www.bd.com



EC DECLARATION OF CONFORMITY

Manufacturer:	Becton, Dickinson and Company Belliver Industrial Estate, Belliver Way, Roborough, Plymouth, PL6 7BP, United Kingdom Becton, Dickinson and Company Belliver Industrial Estate, Belliver Way, Roborough, Plymouth, PL6 7BP, United Kingdom		
Manufacturing Site(s):			
Products:			
	Catalogue number	Device name	GMDN Code
	365308	BD Vacutainer® LH 68 I.U. Plus Blood Collection Tubes	47589
	367526	BD Vacutainer® LH 170 I.U. Plus Blood Collection Tubes	47589
	367883	BD Vacutainer® LH 68 I.U. Plus Blood Collection Tubes	47589
	367885	BD Vacutainer® LH 102 I.U. Plus Blood Collection Tubes	47589
	368272	BD Vacutainer® LH 34 I.U. Plus Blood Collection Tubes	47589
	368494	BD Vacutainer® LH (Lithium Heparin) 34 I.U. Plus Blood Collection Tubes	47589
	368495	BD Vacutainer® LH 34 I.U. Plus Blood Collection Tubes	47589
	368496	BD Vacutainer® LH 68 I.U. Plus Blood Collection Tubes	47589
	368884	BD Vacutainer® LH (Lithium Heparin) 68 I.U. Plus Blood Collection Tubes	47589
	368886	BD Vacutainer® LH (Lithium Heparin) 102 I.U. Plus Blood Collection Tubes	47589
	368889	BD Vacutainer® LH 102 I.U. Plus Blood Collection Tubes	47589
	369622	BD Vacutainer® NH 102 I.U. Plus Blood Collection Tubes	47589
	367869	BD Vacutainer® NH (Sodium Heparin) 68 I.U. Plus Blood Collection Tubes	47592
	367876	BD Vacutainer® NH (Sodium Heparin) 102 I.U. Plus Blood Collection Tubes	47592
	369623	BD Vacutainer® NH 68 I.U. Plus Blood Collection Tubes	47592
IVDD Classification:	Non Annex II In Vitro Diagnostic Medical Device		
IVDD Conformity Assessment Route:	Annex III (excluding Annex III.6)		

We herewith declare that the above mentioned products meet the provisions of the Council Directive 98/79/EC for *in vitro* diagnostic medical devices. This declaration is based on conformity to Annex III (excluding Annex III.6). All supporting documentation is retained under the premises of the manufacturer.

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List of Harmonized Standards:

EN ISO 13485:2012 Medical devices — Quality management systems — Requirements for regulatory purposes EN ISO 14971:2012 Medical Devices – Application of risk management to medical devices EN 556-1:2001 Sterilisation of medical devices – Requirements for medical devices to be designated "STERILE" – Part 1: Requirements for terminally sterilized medical devices EN ISO 11137-1:2015 Sterilization of health care products – Radiation – Part 1: Requirements for development, validation and routine control of a sterilization process for medical devices EN ISO 11137-2:2015 Sterilization of health care products — Radiation — Part 2: Establishing the sterilization dose. EN ISO 11737-2:2009 Sterilization of medical devices - Microbiological methods Part 2: Tests of sterility performed in the definition, validation and maintenance of a sterilization process EN 14820:2004 Single-use containers for human venous blood specimen collection EN 62366:2008 Medical devices - Application of usability engineering to medical devices EN ISO 18113-1: 2011 In vitro diagnostic medical devices – Information supplied by the manufacturer (Labelling). Part 1: Terms, definitions and general requirements (ISO 18113-1:2009) EN ISO 18113-2: 2011 In vitro diagnostic medical devices – Information supplied by the manufacturer (Labelling). Part 2: In vitro diagnostic reagents for professional use (ISO 18113-2:2009) EN ISO 15223-1:2016 Medical Devices – Symbols to be used with medical device labels, labelling and information to be supplied – Part 1: General Requirements

List of Non-Harmonised Standards:

ISO 14001:2015 Environmental management systems - Requirements with guidance for use EN ISO 11137-3:2017 Sterilisation of health care product – Radiation – part 3: guidance on dosimetric aspects of development, validation and routine control EN ISO 11737-1:2018 Sterilization of medical devices - Microbiological methods - Part 1: Determination of a population of microorganisms on products ISO 6710:1995 Single-Use Containers for Venous Blood Specimen Collection EN ISO 14698-1:2003 Cleanrooms and associated controlled environments -- Biocontamination control -- Part 1: General principles and methods EN ISO 14698-2:2003 Cleanrooms and associated controlled environments -- Biocontamination control -- Part 2: Evaluation and interpretation of biocontamination data EN ISO 14644-1:2015 Cleanrooms and associated controlled environments - Part 1: Classification of air cleanliness EN ISO 14644-2:2015 Cleanrooms and associated controlled environments -- Part 2: Monitoring to provide evidence of cleanroom performance related to air cleanliness by particle concentration ISO 2859-1:1999 Sampling procedures for inspection by attributes - Part Sampling schemes indexed by acceptance quality limit (AQL) for lot-by-lot inspection ASTM D5276:1998 (R 2009) Standard Test Method for Drop Test of Loaded Containers by Free Fall ASTM D999: 2008 (R2015) Standard Test Methods for Vibration Testing of Shipping Containers ASTM D4169: 2014 Standard for Performance Testing of Shipping Containers and Systems ASTM D4728: 2006 (R2012) Standard Test Method for Random Vibration Testing of Shipping Containers ASTM D-775: 1980 (R 1986) Standard Test Method for Drop Test for Loaded Boxes

SIGNED FOR AND ON BEHALF OF:

Becton, Dickinson and Company

PLACE, DATE OF ISSUE:

Plymouth, 18th September 2018

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Brad Spring

Vice President, Regulatory Affairs

BD Life Sciences

Document Number: VR4310006

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	VERSION HISTORY
Current Version Prepared By: Joseph Statham	
REV.	Version Description
Α	Transferred from QDMS to ECC – Version number remained
В	Transfer into new IVD Declaration of Conformity Template (MED-RA-001D)
С	Update EN ISO 11737-1:2006 to EN ISO 11737-1:2018 as per CAPA 325553.