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		
Manufacturing Site(s):	Becton, Dickinson and Company Belliver Industrial Estate, Belliver Way, Roborough, Plymouth, PL6 7BP, United Kingdom		
Products:	Catalogue number	Device name	
	362080	BD Vacutainer® CAT Plus Blood Collection Tubes	
	362081	BD Vacutainer® CAT Plus Blood Collection Tubes	
	362082	BD Vacutainer® CAT Plus Blood Collection Tubes	
	362091	BD Vacutainer® CAT Plus Blood Collection Tubes	
	362092	BD Vacutainer® CAT Plus Blood Collection Tubes	
	364660	BD Vacutainer® CAT Plus Blood Collection Tubes	
	365904	BD Vacutainer® SERUM/CAT Plus Blood Collection Tubes	
	367837	BD Vacutainer® CAT Plus Blood Collection Tubes	
	367895	BD Vacutainer® CAT Plus Blood Collection Tubes	
	367896	BD Vacutainer® CAT (Clot Activator Tube) Plus Blood Collection Tubes	
	368271	BD Vacutainer® SERUM/CAT Plus Blood Collection Tubes	
	368492	BD Vacutainer® CAT (Clot Activator Tube) Plus Blood Collection Tubes	
	368493	BD Vacutainer® CAT Plus Blood Collection Tubes	
	368813	BD Vacutainer® CAT Plus Blood Collection Tubes	
	368814	BD Vacutainer® CAT Plus Blood Collection Tubes	
	368815	BD Vacutainer® CAT (Clot Activator Tube) Plus Blood Collection Tubes	
	368817	BD Vacutainer® CAT Plus Blood Collection Tubes	
	368863	BD Vacutainer® CAT Plus Blood Collection Tubes	
	368975	BD Vacutainer® CAT Plus Blood Collection Tubes	
	369032	BD Vacutainer® CAT (Clot Activator Tube) Plus Blood Collection Tubes	
	369625	BD Vacutainer® CAT Plus Blood Collection Tubes	
VDD Classification:	Non Annex II In Vitro Diagnostic Medical Device		
IVDD Conformity Assessment Route:	Annex III (excluding Annex III.6)		
GMDN:	42386		

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.

## 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-

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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 1: 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

Brad Spring

Vice President, Regulatory Affairs

**BD Life Sciences** 

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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.	