

EC DECLARATION OF CONFORMITY

Legal Manufacturer:	Becton, Dickinson and Company (BD) 1 Becton Drive Franklin Lakes, NJ 07417 USA
Authorized Representative:	Becton Dickinson Ireland Ltd. Donore Road Drogheda Co. Louth A92 YW26 Ireland
Manufacturing Site(s):	<p>Manufacturing: BD Vacutainer® Safety-Lok™ Blood Collection Set Becton, Dickinson and Company (BD) 1575 Airport Road PO Box 2128 Sumter, SC 29153 USA</p> <p>Manufacturing and Sterilization: BD Vacutainer® Safety-Lok™ Blood Collection Set Nipro Medical Industries, Ltd. Tatebayashi Plant 2-19-64 Matsubara, Tatebayashi-shi Gunma, 374-8518 Japan</p> <p>Nipro (Thailand) Corporation Limited 10/2 Moo 8 Bangnomko, Sena Phra Nakhon Si Ayutthaya 13110, Thailand</p>
Products:	<p>362093 BD Vacutainer® Safety-Lok™ Blood Collection Set 21Gx ¾ 362094 BD Vacutainer® Safety-Lok™ Blood Collection Set 23Gx ¾ 362095 BD Vacutainer® Safety-Lok™ Blood Collection Set 25Gx ¾ 367246 BD Vacutainer® Safety-Lok™ Blood Collection Set 21Gx ¾ 367247 BD Vacutainer® Safety-Lok™ Blood Collection Set 23Gx ¾ 367282 BD Vacutainer® Safety-Lok™ Blood Collection Set 21Gx ¾ 367284 BD Vacutainer® Safety-Lok™ Blood Collection Set 23Gx ¾</p>

**TITLE: Declaration of Conformity for
Vacutainer® Brand Safety-Lok™ Blood Collection Set**

	<p>367286 BD Vacutainer® Safety-Lok™ Blood Collection Set 21Gx 3/4</p> <p>367288 BD Vacutainer® Safety-Lok™ Blood Collection Set 23Gx 3/4</p> <p>367295 BD Vacutainer® Safety-Lok™ Blood Collection Set 25Gx 3/4</p> <p>368382 BD Vacutainer® Safety-Lok™ Blood Collection Set 25Gx 3/4</p> <p>368383 BD Vacutainer® Safety-Lok™ Blood Collection Set 25Gx 3/4</p>
Classification:	<p>EU Class IIa Medical Device as defined in the Medical Device Directive 93/42/EEC), Annex IX, Section 2.3, Rule 7: which states that all surgically invasive devices intended for short term use, to which the exceptions do not apply.</p> <p>Canada Class II per Schedule 1, Canadian Medical Device Regulations (CMDR), SOR//98-282 which states that all surgically invasive devices are classified as Class II in which none of the indents apply.</p>
Conformity Assessment Route:	Annex II, Medical Device Directive 93/42/EEC
GMDN:	<p>GMDN Code: 58497</p> <p>GMDN Term: Blood collection set, invasive</p>

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 at the premises of the manufacturer.

Harmonized Standards:	<p>EN 1041:2013</p> <p>EN ISO 10993 - Series</p> <p>EN ISO 11135:2014EN ISO 13485:2016</p> <p>EN 1707:1997</p> <p>EN-ISO-15223-1:2016</p> <p>EN ISO 11607-1:2010</p> <p>EN ISO 11737-1:2018</p> <p>EN ISO 14971:2019</p> <p>EN ISO 14155:2011</p>
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Non-Harmonized Standards	ASTM D5276-98:1998 ASTM D999:2008 ASTM D-4169:2014 ISO 11737-1:2018 AMD 2021
Notified Body:	National Standards Association of Ireland (NSAI) 1 Swift Square Northwood Santry, Dublin 9, Ireland Phone: 353 (01) 807-3800 Fax: 353 (01) 807-3838
CE Certificate Number:	252.191
Date of issuance of original CE certificate:	27 April 1997

Date: [25-Jul-2022](#)

DocuSigned by:
Anne Zavertnik
 Signer Name: Anne Zavertnik
Signing Reason: I approve this document
Signing Time: 25-Jul-2022 | 7:45:18 PM BST
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Anne Zavertnik
Vice President, Regulatory Affairs
Integrated Diagnostic Solutions

<u>REVISION HISTORY</u>		
Current Version Prepared By: Eileen Hiller		
REV.	Revision Description	Releasing ECO (if applicable)
04	Initial Release of new DoC template which incorporates requirements of MED-RA-001C. Previous revision histories are contained in the DoC up to Rev. 03.	N/A
05	Removed EN980:2008 and revised EN ISO 13485:2012 to EN ISO 13485:2016, revised EN ISO 15223-1:2012 to EN ISO 15223-1:2016 in the Harmonized Standards section.	N/A
06	Updated Standards revision dates to comply with V08-510-01.	N/A
07	Updated to “Becton, Dickinson and Company (BD) to align with our certification.	N/A
08	Updated authorized approval to Bradford Spring, VP Regulatory Affairs.	N/A 16-Nov-2018
09	Removed “Blood Collection Sets” from the title as it no longer applies. Changed the Authorized Rep to BD Switzerland Sarl; changed authorized signature to Kay Taylor.	N/A August 2019
10	Update sterilization standard to ISO 11737-1:2018 per BDVS-2020-04-29-085157; updated header to IDS, Specimen Management.	N/A June 2020
11	Update GMDN code to 58497 and GMDN term to align with 58497 code per 252.191.36.	N/A December 2021
12	Corrected GMDN term to align with revised code 58497 per 252.191.36. Added ISO 11737-1:2018 AMD 2021 to non-harmonized standards list per BDVS-2021-12-17-102739	N/A January 2022
13	Updated European Authorized Representative from BD Switzerland to BD Ireland. Change to the EU Authorized Representative name and address due to dissolution of the Swiss-EU mutual recognition agreement. NSAI Regulatory Statement Letter accepting the appointment of BD Ireland as the EAR, dated 24 Feb 2022. Update any references to EN ISO 14971:2012 to: EN ISO 14971:2019 per IDSQUALITYPLAN7591	N/A May 2022
14	Update Nipro Thailand Address, corrected spelling of Ayutthaya, inserted “h” to correct spelling.	N/A July 2022