

EU Technical Documentation Assessment Certificate Medical Device Regulation 2017/745

The National Standards Authority of Ireland (NSAI) as a duly designated Notified Body, (identification number 0050) for the purposes of the European Union under MDR 2017/745

HAS ASSESSED THE TECHNICAL DOCUMENTATION SUBMITTED BY

Becton, Dickinson and Company

1 Becton Drive Franklin Lakes, NJ 07417, USA

Manufacturer SRN: US-MF-000019182

Becton Dickinson Ireland Ltd.

Authorised Representative Name and

Address:

Donore Road County Louth

Drogheda, A92 YW26, Ireland

Scope: BD PosiFlush™ syringes

BD PosiFlush™ XS Syringes are intended to be used FOR FLUSHING ONLY of in-situ peripheral intravenous catheters (PIVCs), peripherally inserted central catheters (PICCs), central venous

catheters (CVCs), and implanted venous access ports.

BD PosiFlush™ XS Syringe is not intended for dry product reconstitution, for medication dilution,

or where intravenous therapy with sodium chloride is indicated.

Intended

Using aseptic technique, BD PosiFlush™ XS Syringe can be used on a sterile field.

Purpose BD PosiFlush™ SP Syringes are intended to be used FOR FLUSHING ONLY of in-situ peripheral intravenous catheters (PIVCs), peripherally inserted central catheters (PICCs), central venous

catheters (CVCs), and implanted venous access ports.

BD PosiFlush™ SP Syringe is not intended for dry product reconstitution, for medication dilution,

or where intravenous therapy with sodium chloride is indicated.

BD PosiFlush™ SP Syringe must not be used on a sterile field.

Conclusion: Technical documentation complies with the requirements of Annex IX, Chapter II (where applicable with Section 5.1) of MDR 2017/745.

Product Certificate Number: 745.008D Re-Issued Date:

First Issue Date: 21 December 2027 Expiry Date: 20 December 2027

Site Certificate Number: MD19.2305

Signed:

Approved by:
Lisa Donlon
European Medical Device

European Medical Device Operations Manager Approved by: Dr Majella Geraghty

European Medical Device Operations Manager

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CONDITIONS AND LIMITATIONS: This certificate remains valid on condition that the Approved Quality Management System is maintained in an adequate and efficacious manner in line with the requirements of the Regulation and shall be subject to surveillance audits carried out by the Notified Body. This certificate is based on examination of identified relevant CS, harmonised standards, test reports and audit reports maintained on file with NSAI. Information on examination and tests as per Annex XII, section 10, is available on request. Changes which could affect conformity with the General Safety and Performance Requirements of MDR 2017/745 or with the conditions prescribed for use of the product must receive further approval from NSAI.

The validity of this certificate depends on conditions and/or is limited to the following:

National Standards Authority of Ireland, 1 Swift Square, Northwood, Santry, Dublin 9, Ireland.

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Appendix I Devices Covered by Certificate					
306573	038290WKCQDZQWJK	BD PosiFlush™ SP Syringes 3ml	III		
306583	038290WKCQDZQWJK	BD PosiFlush™ SP Syringes 3ml EMA/CIS	III		
30657371	038290WKCQDZQWJK	BD PosiFlush™ SP Syringes 3ml India	III		
306574	038290WKCQDZQWJK	BD PosiFlush™ SP Syringes 5ml	III		
306584	038290WKCQDZQWJK	BD PosiFlush™ SP Syringes 5ml EMA/CIS	III		
30657471	038290WKCQDZQWJK	BD PosiFlush™ SP Syringes 5ml India	III		
306575	038290WKCQDZQWJK	BD PosiFlush™ SP Syringes 10ml	III		
306585	038290WKCQDZQWJK	BD PosiFlush™ SP Syringes 10ml EMA/CIS	III		
30657571	038290WKCQDZQWJK	BD PosiFlush™ SP Syringes 10ml India	III		
306570	038290WKCQDZQWJK	BD PosiFlush™ XS Syringes 3ml	III		
306580	038290WKCQDZQWJK	BD PosiFlush™ XS Syringes 3ml EMA/CIS	III		
306571	038290WKCQDZQWJK	BD PosiFlush™ XS Syringes 5ml	III		
306581	038290WKCQDZQWJK	BD PosiFlush™ XS Syringes 5ml EMA/CIS	III		
306572	038290WKCQDZQWJK	BD PosiFlush™ XS Syringes 10ml	III		
306582	038290WKCQDZQWJK	BD PosiFlush™ XS Syringes 10ml EMA/CIS	III		

Appendix II Certificate History				
n/a	n/a	n/a	n/a	