



CERTIFICADO DE EXAMEN CE DE TIPO
de acuerdo con el Anexo III de la Directiva 93/42/CEE

EC TYPE-EXAMINATION CERTIFICATE
in accordance with Annex III of Directive 93/42/EEC

Certificado nº/Certificate no	Fecha de validez/Date of validity	ON nº/NB no
2010 02 0701 ET	Desde/From 25-05-2020 Hasta/To 24-05-2024	0318

A favor de/In favour of:

Fabricante/Manufacturer:

Nombre/Name: BECTON DICKINSON S.A.

Dirección/Address: Camino de Valdeoliva, s/n. 28750- SAN AGUSTÍN DEL GUADALIX (Madrid), España

Representante autorizado ante la UE/Authorized EU representative: Idem

Para el producto/For the product:

Categoría/Category: Productos de un solo uso / *Single-use products*

Grupo genérico/ Instrumentos para punción, inyección y/o extracción de fluidos / *Instruments for*

Generic group: *puncture, injection and/or aspiration of fluids*

Tipo/Type: Especificado en Anexos de este Certificado / *Specified in Annexes to this Certificate*

Elaborado en/In the facilities:

Camino de Valdeoliva, s/n. 28750- SAN AGUSTÍN DEL GUADALIX (Madrid). España
ROAD 31 KM. 24.3 P.O. BOX 4010 JUNCOS, PUERTO RICO 00777-4010

Fecha inicial / Initial date: 08-02-2010

Fecha de prórroga anterior / Previous extension date: 25-05-2015

Este certificado debe ir acompañado por un certificado de alguno de los procedimientos previstos en el art. 11 de la Directiva. / *This certificate must be accompanied by a certificate of one of the procedures foreseen in article 11 of the Directive.* / Este certificado cuya documentación técnica está contenida en el expediente nº 95 04 0005, garantiza que los productos descritos cumplen los requisitos de la Directiva. / *This certificate whose technical documentation is contained in dossier no 95 04 0005, guarantees that the described products fulfils the requirements of the Directive.*

Madrid, 22 de mayo de 2020

DIRECTORA DE LA AGENCIA ESPAÑOLA DE MEDICAMENTOS Y PRODUCTOS SANITARIOS

 **agencia española de
medicamentos y
productos sanitarios**

Fdo. Mª Jesús Lamas Díaz

Agencia Española de Medicamentos y Productos Sanitarios

Fecha de la firma: 22/05/2020

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ORGANISMO NOTIFICADO 0318

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ANEXO N°/ANNEX NO: I

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	25-05-2020	24-05-2024	

A favor de/In favour of:

Fabricante/Manufacturer:

Nombre/Name: BECTON DICKINSON S.A.

Dirección/Address: Camino de Valdeoliva, s/n. 28750- SAN AGUSTÍN DEL GUADALIX (Madrid), España

Representante autorizado ante la UE/Authorized EU representative: Idem

Tipo de producto / Devices type: Agujas espinales / Spinal needles

Clasificación / Classification: III

Código NANDO / NANDO code: MD 0102, MDS 7006

1. Agujas espinales punta tipo Quincke / Spinal needle Quincke type point

1.1. Aguja espinal estéril punta tipo Quincke / Sterile spinal needle Quincke type point: BD Spinal Needle

- 1.1.a 18G × 3.00" (1.2 × 75 mm)
- 1.1.b 18G × 3.50" (1.2 × 90 mm)
- 1.1.c 19G × 3.00" (1.1 × 75 mm)
- 1.1.d 19G × 3.50" (1.1 × 90 mm)
- 1.1.e 20G × 1.50" (0.90 × 38 mm)
- 1.1.f 20G × 3.00" (0.90 × 75 mm)
- 1.1.g 20G × 3.50" (0.90 × 90 mm)
- 1.1.h 22G × 1.50" (0.7 × 38 mm)
- 1.1.i 22G × 2.50" (0.7 × 63 mm)
- 1.1.j 22G × 3.00" (0.7 × 75 mm)
- 1.1.k 22G × 3.50" (0.7 × 90 mm)
- 1.1.l 23G × 3.50" (0.64 × 90 mm)
- 1.1.m 25G × 1.00" (0.50 × 25 mm) NEONATAL
- 1.1.n 25G × 2.00" (0.50 × 51 mm) NEONATAL
- 1.1.o 25G × 3.00" (0.50 × 75 mm)
- 1.1.p 25G × 3.50" (0.50 × 90 mm)
- 1.1.q 26G × 3.50" (0.45 × 90 mm)
- 1.1.r 27G × 3.50" (0.40 × 90 mm)

Agencia Española de Medicamentos y Productos Sanitarios

Fecha de la firma: 22/05/2020

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1.2. Aguja espinal no estéril punta tipo Quincke / Non-sterile spinal needle Quincke type point: BD Spinal Needle

1.2.a 22G × 3.50" (0.70 × 90 mm)

1.2.b 25G × 3.50" (0.50 × 90 mm)

1.2.c 26G × 3.50" (0.45 × 90 mm)

1.3. Set aguja espinal punta tipo Quincke con introductor, estéril / Sterile spinal needle Quincke type point with introducer set: BD Spinal Needle

1.3.a 25G × 3.50" (0.50 × 90 mm) con introductor / with introducer 20G × 1.25" (0.9 × 32 mm)

1.3.b 26G × 3.50" (0.45 × 90 mm) con introductor / with introducer 20G × 1.25" (0.9 × 32 mm)

1.3.c 27G × 3.50" (0.40 × 90 mm) con introductor / with introducer 20G × 1.25" (0.9 × 32 mm)

Este certificado ampara todas las marcas de estos productos incluidas por el fabricante en su declaración CE de conformidad. This certificate covers all trademarks of these products included by the manufacturer in his EC declaration of conformity.

Madrid, 22 de mayo de 2020

DIRECTORA DE LA AGENCIA ESPAÑOLA DE MEDICAMENTOS Y PRODUCTOS SANITARIOS

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Fdo. Mª Jesús Lamas Díaz

Agencia Española de Medicamentos y Productos Sanitarios

Fecha de la firma: 22/05/2020

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BDCERTIFICADO CE DE GARANTÍA DE CALIDAD DE LA PRODUCCIÓN
de acuerdo con el Anexo V de la Directiva 93/42/CEE
EC PRODUCTION QUALITY ASSURANCE CERTIFICATE
in accordance with Annex V of Directive 93/42/EEC

Certificado nº/Certificate no	Fecha de validez/Date of validity	ON nº/NB no
95 06 0005 CP	Desde/From 26-05-2021 Hasta/To 24-05-2024	0318

A favor de/In favour of:

Fabricante/Manufacturer:

Nombre/Name: BECTON DICKINSON S.A.

Dirección/Address: Camino de Valdeoliva, s/n. 28750- SAN AGUSTÍN DEL GUADALIX (Madrid),
España

Representante autorizado ante la UE / Authorized EU representative: Idem

Para el producto/For the product:

Categoría/Category: Productos de un solo uso / *Single-use products*

Grupo genérico/ Instrumentos para punción, inyección y/o extracción de fluidos / *Instruments for*
Generic group: *puncture, injection and/or aspiration of fluids*

Tipo/Type: Especificado en Anexos de este Certificado / *Specified in Annexes to this Certificate*

Elaborado en/In the facilities:

Camino de Valdeoliva, s/n. 28750- SAN AGUSTÍN DEL GUADALIX (Madrid). España
ROAD 31 KM. 24.3 P.O. BOX 4010 JUNCOS, PUERTO RICO 00777-4010

Fecha inicial / Initial date: 19-06-1995

Fecha de prórroga anterior / Previous extension date: 25-05-2020

Este certificado debe ir acompañado por / *This certificate must be accompanied by:* Certificado de examen CE de tipo (Anexo III) para los productos de clase III y Declaración CE de conformidad (anexo VII) para los productos de clase IIa / *EC Type examination certificate (Annex III) for Class III medical devices and EC Declaration of conformity (Annex VII) for class IIa medical devices.*

Este certificado es consecuencia de la auditoria del sistema completo de garantía de calidad de la producción y del examen de la documentación técnica contenida en el expediente nº 95 04 0005, y garantiza que los productos descritos cumplen los requisitos de la Directiva. / *This certificate is issued on the production quality assurance system audit, and the examination of the technical documentation contained in dossier no 95 04 0005, and guarantees that the described products fulfils the requirements of the Directive.*

Madrid, 25 de mayo de 2021

DIRECTORA DE LA AGENCIA ESPAÑOLA DE MEDICAMENTOS Y PRODUCTOS SANITARIOS



agencia española de
medicamentos y
productos sanitarios

Fdo. Mª Jesús Lamas Díaz

Agencia Española de Medicamentos y Productos Sanitarios (AEMPS)

Fecha de la firma: 25/05/2021

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ORGANISMO NOTIFICADO 0318



ANEXO N°/ANNEX NO: I

CERTIFICADO CE DE GARANTÍA DE CALIDAD DE LA PRODUCCIÓN
de acuerdo con el Anexo V de la Directiva 93/42/CEE

*EC PRODUCTION QUALITY ASSURANCE CERTIFICATE
in accordance with Annex V of Directive 93/42/EEC*

Certificado n°/Certificate no	Fecha de validez/Date of validity		ON n°/NB no
95 06 0005 CP	Desde/From	Hasta/To	0318
	26-05-2021	24-05-2024	

A favor de/In favor of:

Fabricante/Manufacturer:

Nombre/Name: BECTON DICKINSON S.A.

Dirección/Address: Camino de Valdeoliva, s/n. 28750- SAN AGUSTÍN DEL GUADALIX (Madrid), España

Representante autorizado ante la UE / Authorized EU representative: Idem

Tipo de producto / Devices type: Aguja espinal, introductor de agujas espinales, jeringas con y sin aguja, jeringas para insulina / Spinal needle, needle introducer, syringes with and without needle and insulin syringes.

Clasificación / Classification: III

1 Aguja espinal / Spinal needles

Código NANDO / NANDO code: MD 0102

1.1 Aguja espinal Whitacre punta tipo lápiz / Whitacre Pencil Point Spinal Needles (Descrito en el certificado de examen CE de tipo n° 2010 02 0700 ET / Described in the EC Type-examination certificate no 2010 02 0700 ET)

1.1.a Aguja espinal Whitacre punta tipo lápiz estéril / Sterile Whitacre Pencil Point Spinal Needle: BD Whitacre Needle

1.1.a.1. 22G × 3.50" (0.7 × 90 mm)

1.1.a.2. 24G × 3.50" (0.55 × 90 mm)

1.1.a.3. 25G × 3.50" (0.50 × 90 mm)

1.1.a.4. 25G × 4.06" (0.50 × 103 mm)

1.1.a.5. 27G × 3.50" (0.40 × 90 mm)

1.1.a.6. 27G × 4.06" (0.40 × 103 mm)

Agencia Española de Medicamentos y Productos Sanitarios (AEMPS)

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1.1.b Set aguja espinal Whitacre punta tipo lápiz con introductor, estéril / *Sterile Whitacre Pencil Point Spinal Needle with introducer set*: **BD Whitacre Needle**

1.1.b.1. 25G × 3.50" (0.50 × 90 mm) con introductor / *with introducer* 20G × 1.25" (0.9 × 32 mm)

1.1.b.2. 25G × 4.06" (0.50 × 103 mm) con introductor / *with introducer* 20G × 1.25" (0.9 × 32 mm)

1.1.b.3. 27G × 3.50" (0.40 × 90 mm) con introductor / *with introducer* 22G × 1.25" (0.9 × 32 mm)

1.1.b.4. 27G × 4.06" (0.40 × 103 mm) con introductor / *with introducer* 22G × 1.25" (0.9 × 32 mm)

1.2. Agujas espinales punta tipo Quincke / Spinal needle Quincke type point (Descrito en el certificado de examen CE de tipo n° 2010 02 0701 ET / Described in the EC Type-examination certificate no 2010 02 0701 ET)

1.2.a Aguja espinal estéril punta tipo Quincke / *Sterile spinal needle Quincke type point*: **BD Spinal Needle**

1.2.a.1. 18G × 3.00" (1.2 × 75 mm)

1.2.a.2. 18G × 3.50" (1.2 × 90 mm)

1.2.a.3. 19G × 3.00" (1.1 × 75 mm)

1.2.a.4. 19G × 3.50" (1.1 × 90 mm)

1.2.a.5. 20G × 1.50" (0.90 × 38 mm)

1.2.a.6. 20G × 3.00" (0.90 × 75 mm)

1.2.a.7. 20G × 3.50" (0.90 × 90 mm)

1.2.a.8. 22G × 1.50" (0.7 × 38 mm)

1.2.a.9. 22G × 2.50" (0.7 × 63 mm)

1.2.a.10. 22G × 3.00" (0.7 × 75 mm)

1.2.a.11. 22G × 3.50" (0.7 × 90 mm)

1.2.a.12. 23G × 3.50" (0.64 × 90 mm)

1.2.a.13. 25G × 1.00" (0.50 × 25 mm) Neonatal

1.2.a.14. 25G × 2.00" (0.50 × 51 mm) Neonatal

Agencia Española de Medicamentos y Productos Sanitarios (AEMPS)

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1.2.a.15. 25G × 3.00" (0.50 × 75 mm)

1.2.a.16. 25G × 3.50" (0.50 × 90 mm)

1.2.a.17. 26G × 3.50" (0.45 × 90 mm)

1.2.a.18. 27G × 3.50" (0.40 × 90 mm)

1.2.b Aguja espinal no estéril punta tipo Quincke / *Non-sterile spinal needle Quincke type point: BD Spinal Needle*

1.2.b.1. 22G × 3.50" (0.70 × 90 mm)

1.2.b.2. 25G × 3.50" (0.50 × 90 mm)

1.2.b.3. 26G × 3.50" (0.45 × 90 mm)

1.2.c Set aguja espinal punta tipo Quincke con introductor, estéril / *Sterile spinal needle Quincke type point with introducer set: BD Spinal Needle*

1.2.c.1. 25G × 3.50" (0.50 × 90 mm) con introductor / *with introducer* 20G × 1.25" (0.9 × 32 mm)

1.2.c.2. 26G × 3.50" (0.45 × 90 mm) con introductor / *with introducer* 22G × 1.25" (0.9 × 32 mm)

1.2.c.3. 27G × 3.50" (0.40 × 90 mm) con introductor / *with introducer* 22G × 1.25" (0.9 × 32 mm)

1.3 Aguja espinal BD Whitacre NRFit™ / *BD Whitacre Spinal NRFit™ Needles*

(Descrito en el certificado de examen CE de tipo n° 2020 04 0912 ET / Described in the EC Type-examination certificate no 2020 04 0912 ET)

1.3.a Aguja espinal BD Whitacre NRFit™ estéril / *Sterile BD Whitacre Spinal NRFit™ Needle BD Whitacre Spinal NRFit™ Needle*

1.3.a.1 25G × 4.06" (0.50 × 103.2 mm)

1.3.a.2 25G × 4.70" (0.50 × 119.1 mm)

1.3.a.3 27G × 4.06" (0.40 × 103.2 mm)

Agencia Española de Medicamentos y Productos Sanitarios (AEMPS)

Fecha de la firma: 25/05/2021

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1.3.a.4	27G × 4.70" (0.40 × 119.1 mm)
1.3.a.5	22G × 3.50" (0.70 × 88.9 mm)
1.3.a.6	24G × 3.50" (0.55 × 88.9 mm)
1.3.a.7	25G × 3.50" (0.50 × 88.9 mm)
1.3.a.8	27G × 3.50" (0.40 × 88.9 mm)
1.3.a.9	20G × 3.5" (0.90 × 88.9 mm)
1.3.a.10	21G × 3.5" (0.80 × 88.9 mm)
1.3.a.11	22G × 4" (0.70 × 101.6 mm)
1.3.a.12	26G × 3.5" (0.45 × 88.9 mm)

1.3.b Agujas espinales BD Whitacre NRFit™ no estériles / *Non-sterile BD Whitacre Spinal NRFit™ Needles*
BD Whitacre Spinal NRFit™ Needle

1.3.b.1	25G × 4.06" (0.50 × 103.2 mm)
1.3.b.2	27G × 4.06" (0.40 × 103.2 mm)
1.3.b.3	22G × 3.50" (0.70 × 88.9 mm)
1.3.b.4	24G × 3.50" (0.55 × 88.9 mm)
1.3.b.5	25G × 3.50" (0.50 × 88.9 mm)
1.3.b.6	25G × 4.70" (0.50 × 119.1 mm)
1.3.b.7	27G × 3.50" (0.40 × 88.9 mm)
1.3.b.8	27G × 4.70" (0.40 × 119.1 mm)

Agencia Española de Medicamentos y Productos Sanitarios (AEMPS)

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				0318

1.3.c Set aguja espinal BD Whitacre NRFit™ con introductor, estéril / *Sterile BD Whitacre Spinal NRFit™ Needle with introducer set. BD Whitacre Spinal NRFit™ Needle Set.*

1.3.c.1 27G × 3.50" (0.40 × 88.9 mm) con introductor / *with introducer* 22G × 1.25" (0.70 × 31.8 mm)

1.3.c.2 25G × 3.50" (0.50 × 88.9 mm) con introductor / *with introducer* 20G × 1.25" (0.90 × 31.8 mm)

1.3.c.3 25G × 4.06" (0.50 × 103.2 mm) con introductor / *with introducer* 20G × 1.25" (0.90 × 31.8 mm)

1.3.c.4 27G × 4.06" (0.40 × 103.2 mm) con introductor / *with introducer* 22G × 1.25" (0.70 × 31.8 mm)

1.4 Aguja espinal BD Quincke NRFit™ / BD Quincke Spinal NRFit™ Needles

(Descrito en el certificado de examen CE de tipo n° 2020 04 0913 ET / *Described in the EC Type-examination certificate no 2020 04 0913 ET*)

1.4.a Aguja espinal BD Quincke NRFit™ estéril / *Sterile BD Quincke Spinal NRFit™ Needle* **BD Quincke Spinal NRFit™ Needle**

1.4.a.1 25G × 3.00" (0.5 × 76.2 mm)

1.4.a.2 18G × 3.00" (1.2 × 76.2 mm)

1.4.a.3 18G × 3.50" (1.2 × 88.9 mm)

1.4.a.4 22G × 1.50" (0.7 × 38.1 mm)

1.4.a.5 22G × 2.50" (0.7 × 63.5 mm)

1.4.a.6 22G × 3.00" (0.7 × 76.2 mm)

1.4.a.7 22G × 3.50" (0.7 × 88.9 mm)

1.4.a.8 25G × 1.00" (0.5 × 25.4 mm)

1.4.a.9 25G × 3.50" (0.5 × 88.9 mm)

1.4.a.10 18G × 6.00" (1.20 × 152.4 mm)

1.4.a.11 20G × 6.00" (0.90 × 152.4 mm)

1.4.a.12 22G × 5.00" (0.70 × 127 mm)

Agencia Española de Medicamentos y Productos Sanitarios (AEMPS)

Fecha de la firma: 25/05/2021

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ANEXO N°/ANNEX NO: I

CERTIFICADO CE DE GARANTÍA DE CALIDAD DE LA PRODUCCIÓN
de acuerdo con el Anexo V de la Directiva 93/42/CEE

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95 06 0005 CP	Desde/From	26-05-2021	Hasta/To	24-05-2024
				0318

1.4.a.13 22G × 7.00" (0.70 × 177.8 mm)

1.4.a.14 25G × 4.70" (0.50 × 119.1 mm)

1.4.a.15 25G × 2.00" (0.5 × 50.8 mm)

1.4.a.16 19G × 3.00" (1.1 × 76.2 mm)

1.4.a.17 19G × 3.50" (1.1 × 88.9 mm)

1.4.a.18 20G × 1.50" (0.9 × 38.1 mm)

1.4.a.19 20G × 3.00" (0.9 × 76.2 mm)

1.4.a.20 20G × 3.50" (0.9 × 88.9 mm)

1.4.a.21 23G × 3.50" (0.6 × 88.9 mm)

1.4.a.22 26G × 3.50" (0.45 × 88.9 mm)

1.4.a.23 27G × 3.50" (0.4 × 88.9 mm)

1.4.b Aguja espinal BD Quincke NRFit™ no estériles / *Non-sterile BD Quincke Spinal NRFit™ Needle*
BD Quincke Spinal NRFit™ Needle

1.4.b.1 22G × 3.50" (0.70 × 88.9 mm)

1.4.b.2 22G × 2.50" (0.70 × 63.5 mm)

1.4.b.3 22G × 7.00" (0.70 × 177.8 mm)

1.4.b.4 18G × 3.50" (1.2 × 88.9 mm)

1.4.b.5 22G × 1.50" (0.7 × 38.1 mm)

1.4.b.6 22G × 5.00" (0.70 × 127 mm)

1.4.b.7 25G × 2.00" (0.5 × 50.8 mm)

1.4.b.8 25G × 3.50" (0.5 × 88.9 mm)

1.4.b.9 20G × 3.50" (0.9 × 88.9 mm)

1.4.b.10 26G × 3.50" (0.45 × 88.9 mm)

1.4.b.11 27G × 3.50" (0.4 × 88.9 mm)

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1.4.c Set Aguja espinal BD Quincke NRFit™ con introductor, estéril / *Sterile BD Quincke Spinal NRFit™ Needle with introducer set* **BD Quincke Spinal NRFit™ Needle Set**

1.4.c.1 25G × 3.50" (0.50 × 88.9 mm) con introductor / *with introducer* 20G × 1.25" (0.90 × 31.8 mm)

1.4.c.2 26G × 3.50" (0.45 × 88.9 mm) con introductor / *with introducer* 22G × 1.25" (0.70 × 31.8 mm)

1.4.c.3 27G × 3.50" (0.40 × 88.9 mm) con introductor / *with introducer* 22G × 1.25" (0.70 × 31.8 mm)

Clasificación / Classification: IIa

2. Introductor de agujas espinales / *Needle introducer*

Código NANDO / NANDO code: MD 0102

2.1. BD Introductor de agujas espinales, estéril / *Sterile BD Spinal needle introducer.* **BD Spinal Needle**

2.1.a. 20G × 1.25" (0.90 × 32 mm)

2.1.b. 22G × 1.25" (0.70 × 32 mm)

2.2 BD Aguja introductora espinal NRFit™ estéril / *Sterile BD Spinal needle introducer NRFit™* **BD Spinal needle introducer NRFit™**

2.2.a 20G × 1.25" (0.90 × 31.8 mm)

2.2.b 22G × 1.25" (0.70 × 31.8 mm)

2.3 **BD Aguja introductora espinal NRFit™ no estéril / *Non-sterile BD Spinal needle introducer NRFit™*** **BD Spinal needle introducer NRFit™**

2.3.a 20G × 1.25" (0.90 × 31.8 mm)

2.3.b 22G × 1.25" (0.70 × 31.8 mm)

2.3.c. 18G × 1.25" (1.2 × 31.8 mm)

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3. Jeringas de tres piezas con aguja / Three-piece syringes with needle

Código NANDO / NANDO code: MD 0102

3.1. Jeringas estériles BD Plastipak™ cono Luer con aguja BD Microlance™ 3 / Sterile BD Plastipak™ Luer-Slip syringes with BD Microlance™ 3 needle

3.1.a. Jeringas de: / Syringes of: 1 ml

- 3.1.a.1. Con aguja / With needle: 25G × 5/8" (0.5 × 16 mm)
- 3.1.a.2. Con aguja / With needle: 26G × 3/8" (0.45 × 10 mm)
- 3.1.a.3. Con aguja / With needle: 23G × 1" (0.6 × 25 mm)
- 3.1.a.4. Con aguja / With needle: 22G × 1 1/2" (0.7 × 40 mm)
- 3.1.a.5. Con aguja / With needle: 22G × 1 1/4" (0.7 × 30 mm)
- 3.1.a.6. Con aguja / With needle: 27G × 3/8" (0.4 × 10 mm)

3.1.b. Jeringas de: / Syringes of: 2 ml

- 3.1.b.1. Con aguja / With needle: 21G × 5/8" (0.8 × 16 mm)
- 3.1.b.2. Con aguja / With needle: 21G × 1 1/2" (0.8 × 40 mm)
- 3.1.b.3. Con aguja / With needle: 22G × 1 1/4" (0.7 × 30 mm)
- 3.1.b.4. Con aguja / With needle: 23G × 1" (0.6 × 25 mm)
- 3.1.b.5. Con aguja / With needle: 25G × 5/8" (0.5 × 16 mm)

3.1.c. Jeringas de: / Syringes of: 5 ml

- 3.1.c.1. Con aguja / With needle: 21G × 1 1/2" (0.8 × 40 mm)
- 3.1.c.2. Con aguja / With needle: 22G × 1 1/4" (0.7 × 30 mm)

3.1.d. Jeringas de: / Syringes of: 10 ml

- 3.1.d.1. Con aguja / With needle: 21G × 1 1/2" (0.8 × 40 mm)

3.1.e. Jeringas de: / Syringes of: 20 ml

- 3.1.e.1. Con aguja / With needle: 21G × 1 1/2" (0.8 × 40 mm)

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3.2. Jeringas estériles BD Plastipak™ Luer-Lok™ con aguja BD Microlance™ 3 / Sterile BD Plastipak™ Luer-Lok™ syringes with BD Microlance™ 3 needles

3.2.a. Jeringa de 50 ml con aguja 14G × 1¼" (2.1 × 30 mm) / Syringes of 50 ml with 14G × 1¼" (2.1 × 30 mm) needle

3.2.b. Jeringa de 50 ml color ámbar con aguja 14G × 1¼" (2.1 × 30 mm) / Amber color syringes of 50 ml with 14G × 1¼" (2.1 × 30 mm) needle

3.3. Jeringas estériles BD Plastipak™ Luer-Lok™ con aguja Blunt / Sterile BD Plastipak™ Luer-Lok™ syringes with Blunt Fill Needle

3.3.a. Jeringas de 50 ml con aguja 18G × 1½" (1.2 × 40 mm) / Syringes of 50 ml with needle 18G × 1½" (1.2 × 40 mm)

3.3.b. Jeringas de 50 ml color ámbar con aguja 18G × 1½" (1.2 × 40 mm) / Amber color syringes of 50 ml with needle 18G × 1½" (1.2 × 40 mm)

3.3.c. Jeringas de 50 ml con aguja 18G × 1" (1.2 × 25 mm) / Syringes of 50 ml with Needle 18G × 1" (1.2 × 25 mm)

3.3.d. Jeringas de 50 ml color ámbar con aguja 18G × 1" (1.2 × 25 mm) / Amber color syringes of 50 ml with Needle 18G × 1" (1.2 × 25 mm)

3.4. Jeringas estériles BD Perfusion con aguja BD Microlance™ 3 / Sterile BD Perfusion syringes with BD Microlance™ 3 needle

3.4.a. Jeringas con aguja 14G × 1¼" (2.1 × 30 mm) / Syringes with 14G × 1¼" (2.1 × 30 mm) needle

3.4.b. Jeringas color ámbar con aguja 14G × 1¼" (2.1 × 30 mm) / Amber color syringes with 14G × 1¼" (2.1 × 30 mm) needle

Agencia Española de Medicamentos y Productos Sanitarios (AEMPS)

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4. Jeringas de tres piezas sin aguja / Three-piece syringes without needle

Código NANDO / NANDO code:MD 0102

4.1. Jeringas estériles BD Perfusion / Sterile BD Perfusion syringes

4.1.a. Jeringas de / Syringes of: 50 ml

4.1.b. Jeringas color ambar de / Amber color syringes of: 50 ml

4.1.c. Envase Convenience de jeringas de / Convenience Pack of syringes of: 50 ml

4.2. Jeringas estériles BD Plastipak™ Luer-Lok™ / Sterile BD Plastipak™ Luer-Lok™ syringes

4.2.a. Jeringas de: / Syringes of: 10 ml

4.2.b. Jeringas de: / Syringes of: 20 ml

4.2.c. Envase Convenience de jeringas de: / Convenience Pack of syringes of: 20 ml

4.2.d. Jeringas de: / Syringes of: 30 ml

4.2.e. Jeringas de: / Syringes of: 50 ml

4.2.f. Jeringas de color ámbar: / Amber color syringes of: 50 ml

4.2.g. Envase Convenience de jeringas de: / Convenience Pack of syringes of: 50 ml

4.3. Jeringas no estériles BD Plastipak™ Luer-Lok™ / Non-sterile BD Plastipak™ Luer-Lok™ syringes

4.3.a. Jeringas de: / Syringes of: 20 ml

4.3.b. Jeringas de: / Syringes of: 50 ml





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5. Jeringas de tres piezas cono Luer con aguja para insulina / Three-piece insulin Luer-Slip syringes with needle

Código NANDO / NANDO code: MD 0102

5.1. Jeringas estériles BD Plastipak™ para insulina de 1 ml 40 UI / Sterile BD Plastipak™ Insulin 1ml 40 I.U. syringes

5.1.a.1. Con aguja / With needle: 30G × ½" (0.3 × 13 mm)

5.2. Jeringas estériles BD Plastipak™ para insulina de 1 ml 100 UI / Sterile BD Plastipak™ Insulin 1ml 100 I.U. syringes

5.2.a.1. Con aguja / With needle: 25G × ⅝" (0.5 × 16 mm)

5.2.a.2. Con aguja / With needle: 26G × ⅜" (0.45 × 10 mm)

Este certificado ampara todas las marcas de estos productos incluidas por el fabricante en su declaración CE de conformidad.

This certificate covers all trademarks of these products included by the manufacturer in his EC declaration of conformity.

Madrid, 25 de mayo de 2021

DIRECTORA DE LA AGENCIA ESPAÑOLA DE MEDICAMENTOS Y PRODUCTOS SANITARIOS

 **agencia española de
medicamentos y
productos sanitarios**

Fdo. Mª Jesús Lamas Díaz

Agencia Española de Medicamentos y Productos Sanitarios (AEMPS)

Fecha de la firma: 25/05/2021

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S/REF CNCps ID: 92261
N/REF ON0318/GH/JC/PS/95 04 0005
FECHA: 16 de noviembre de 2021
ASUNTO: FE DE ERRATAS
Certificado CE Nº 95 06 0005 CP
(localizador TJ4EL6PA55)

BECTON DICKINSON S.A.
Camino de Valdeoliva s/n
28750 San Agustín de Guadalix
(MADRID)

Vista la solicitud de la empresa BECTON DICKINSON S.A de corrección de varias erratas en las medidas de los introductores de los diferentes sets de agujas recogidos en los epígrafes 1.1.b.3, 1.1.b.4, 1.2.c.2, 1.2.c.3 y 1.4.c.2 del certificado indicado en el asunto con fecha de entrada en vigor 26.05.2021, se confirma la existencia de dichas erratas y a los efectos oportunos se certifica que:

Donde dice:

- 1.1.b.3.** 27G × 3.50" (0.40 × 90 mm) con introductor / *with introducer* **20G × 1.25" (0.9 × 32 mm)**
- 1.1.b.4.** 27G × 4.06" (0.40 × 103 mm) con introductor / *with introducer* **20G × 1.25" (0.9 × 32 mm)**
- 1.2.c.2.** 26G × 3.50" (0.45 × 90 mm) con introductor / *with introducer* **22G × 1.25" (0.7 × 32 mm)**
- 1.2.c.3.** 27G × 3.50" (0.40 × 90 mm) con introductor / *with introducer* **20G × 1.25" (0.9 × 32 mm)**
- 1.4.c.2.** 26G × 3.50" (0.45 × 88.9 mm) con introductor / *with introducer* **22G × 1.25" (0.7 × 31.8 mm)**

Debe decir:

- 1.1.b.3.** 27G × 3.50" (0.40 × 90 mm) con introductor / *with introducer* **22G × 1.25" (0.7 × 32 mm)**
- 1.1.b.4.** 27G × 4.06" (0.40 × 103 mm) con introductor / *with introducer* **22G × 1.25" (0.7 × 32 mm)**
- 1.2.c.2.** 26G × 3.50" (0.45 × 90 mm) con introductor / *with introducer* **20G × 1.25" (0.9 × 32 mm)**
- 1.2.c.3.** 27G × 3.50" (0.40 × 90 mm) con introductor / *with introducer* **22G × 1.25" (0.7 × 32 mm)**
- 1.4.c.2.** 26G × 3.50" (0.45 × 88.9 mm) con introductor / *with introducer* **20G × 1.25" (0.9 × 31.8 mm)**





Taking account of the BECTON DICKINSON SA request to correct several errors in the introducers measurements of different needle sets included in sections 1.1.b.3, 1.1.b.4, 1.2.c.2, 1.2.c.3 and 1.4.c.2 of the subject certificate with effective date 26.05.2021, it is confirmed the existence of the mentioned errors and for the appropriate purposes, it is certified that:

Where it says:

1.1.b.3. 27G x 3.50" (0.40 x 90 mm) con introductor / with introducer **20G x 1.25" (0.9 x 32 mm)**

1.1.b.4. 27G x 4.06" (0.40 x 103 mm) con introductor / with introducer **20G x 1.25" (0.9 x 32 mm)**

1.2.c.2. 26G x 3.50" (0.45 x 90 mm) con introductor / with introducer **22G x 1.25" (0.7 x 32 mm)**

1.2.c.3. 27G x 3.50" (0.40 x 90 mm) con introductor / with introducer **20G x 1.25" (0.9 x 32 mm)**

1.4.c.2. 26G x 3.50" (0.45 x 88.9 mm) con introductor / with introducer **22G x 1.25" (0.7 x 31.8 mm)**

It should read:

1.1.b.3. 27G x 3.50" (0.40 x 90 mm) con introductor / with introducer **22G x 1.25" (0.7 x 32 mm)**

1.1.b.4. 27G x 4.06" (0.40 x 103 mm) con introductor / with introducer **22G x 1.25" (0.7 x 32 mm)**

1.2.c.2. 26G x 3.50" (0.45 x 90 mm) con introductor / with introducer **20G x 1.25" (0.9 x 32 mm)**

1.2.c.3. 27G x 3.50" (0.40 x 90 mm) con introductor / with introducer **22G x 1.25" (0.7 x 32 mm)**

1.4.c.2. 26G x 3.50" (0.45 x 88.9 mm) con introductor / with introducer **20G x 1.25" (0.9 x 31.8 mm)**

Madrid, 18 de noviembre de 2021

DIRECTORA DE LA AGENCIA ESPAÑOLA DE MEDICAMENTOS Y PRODUCTOS SANITARIOS

 **agencia española de
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productos sanitarios**

Fdo. Mª Jesús Lamas Díaz

Agencia Española de Medicamentos y Productos Sanitarios (AEMPS)

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Supplementary information to AR120 818297

Non-significant changes approved after the 26th May 2021 as per the Transitional Provisions of MDR Article 120.3

Issued to:

BECTON DICKINSON, S.A.

Camino de Valdeoliva, s/n. 28750- SAN AGUSTÍN DEL GUADALIX (Madrid),

España

Date: 10 January 2025

Changes Approved:

Date	Reference Number	Action
10 January 2025	30288755	Transfer of appropriate surveillance to BSI per Regulation (EU) 2023/607 of Sterile BD Plastipak™ Luer-Lok™ Syringe with Blunt Fill Needle (colourless and Amber), Sterile BD Plastipak™ Luer-Lok™ Syringe (without needle), Sterile BD Plastipak™ Luer-Lok™ Convenience Pack (without needle), BD Plastipak™ BD Luer-Lok™ Syringe (without needle Bulk Non-Sterile), Sterile BD Plastipak™ Luer-Slip Syringe with Microlance needle, Sterile BD Plastipak™ Luer-Slip insulin syringe (with needle), Sterile BD Spinal Introducer NRFit™ Needle, BD Spinal Introducer NRFit™ Needle (Bulk Non-Sterile), Sterile BD Whitacre Spinal Needle (see AR120 818313 for variants), Sterile BD Quincke Spinal Needle (see AR120 818312 for variants), Sterile BD Quincke Spinal Needle Set (Spinal Needle + introducer) (see AR120 818312 for variants), Sterile BD Whitacre Spinal Needle Set

Date	Reference Number	Action
10 January 2025	30288755	(Spinal Needle + introducer) (see AR120 818313 for variants), Sterile BD Quincke Spinal NRFit™ Needle (see AR120 818306 for variants), BD Quincke Spinal NRFit™ Needle (Bulk Non-Sterile) (see AR120 818306 for variants), Sterile BD Whitacre Spinal NRFit™ Needle (see AR120 818310 for variants), BD Whitacre Spinal NRFit™ Needle (Bulk Non - Sterile) (see AR120 818310 for variants), Sterile BD Quincke Spinal NRFit™ Needle Set (NRFit Spinal Needle + introducer) (see AR120 818306 for variants), Sterile BD Whitacre Spinal NRFit™ Needle Set (NRFit Spinal Needle + introducer) (see AR120 818310 for variants). Original NB Certificate Number: 95 06 0005 CP

10 January 2025

Becton Dickinson, S.A.
Camino de Valdeoliva, s/n
San Agustin del Guadalix Madrid
28750
Spain

To whom it may concern,

The transitional provisions specified in MDR Article 120(3) (as amended by (EU) 2023/607) prohibit Notified Bodies from issuing new certificates or amending, modifying, supplementing any existing MDD/AIMDD certificates from 26th May 2021.

This letter is to confirm that BSI has reviewed and approved the change(s) detailed in the table below. These changes do not represent a significant change in design or intended purpose under MDR Article 120(3) (as amended by (EU) 2023/607) and as per the guidance provided in MDCG 2020-3.

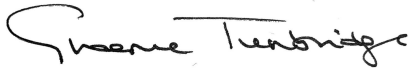
The related MDD certificate specified below remains valid until the expiry date stated on the certificate or until the end of the transition period as specified in Article 120(3c) of MDR (as amended by (EU) 2023/607), subject to the manufacturer's continued compliance to the other conditions provided in Article 120(3c) of MDR (as amended by (EU) 2023/607).

Original Certificate Number	BSI Reference Number	Directive and Annex	Reference Number	Changes approved
95 06 0005 CP	AR120 818297	93/42/EEC Annex V	30288755	Transfer of appropriate surveillance to BSI per Regulation (EU) 2023/607 of of Sterile BD Plastipak™ Luer-Lok™ Syringe with Blunt Fill Needle (colourless and Amber), Sterile BD Plastipak™ Luer-Lok™ Syringe (without needle), Sterile BD Plastipak™ Luer-Lok™ Convenience Pack (without needle), BD Plastipak™ BD Luer-Lok™ Syringe (without needle Bulk Non-Sterile), Sterile BD Plastipak™ Luer-Slip Syringe with Microlance needle, Sterile BD Plastipak™ Luer-Slip insulin syringe (with needle), Sterile BD Spinal Introducer NRFit™ Needle, BD Spinal Introducer NRFit™ Needle (Bulk Non-Sterile), Sterile BD Whitacre Spinal Needle (see AR120 818313 for variants), Sterile BD Quincke Spinal Needle (see AR120 818312 for variants), Sterile BD Quincke Spinal Needle Set (Spinal Needle + introducer) (see AR120 818312 for variants), Sterile BD Whitacre Spinal Needle Set (Spinal Needle + introducer) (see AR120 818313 for variants), Sterile BD Quincke Spinal NRFit™ Needle (see AR120 818306 for variants), BD Quincke Spinal NRFit™ Needle (Bulk Non-

Original Certificate Number	BSI Reference Number	Directive and Annex	Reference Number	Changes approved
95 06 0005 CP	AR120 818297	93/42/EEC Annex V	30288755	Sterile) (see AR120 818306 for variants), Sterile BD Whitacre Spinal NRFit™ Needle (see AR120 818310 for variants), BD Whitacre Spinal NRFit™ Needle (Bulk Non - Sterile) (see AR120 818310 for variants), Sterile BD Quincke Spinal NRFit™ Needle Set (NRFit Spinal Needle + introducer) (see AR120 818306 for variants), Sterile BD Whitacre Spinal NRFit™ Needle Set (NRFit Spinal Needle + introducer) (see AR120 818310 for variants).

Should you have any queries concerning your certification, or if we can be of further assistance to you, please contact your BSI Scheme Manager.

Yours sincerely,



Graeme Tunbridge
Senior Vice President, Medical Devices

Becton Dickinson S.A.
Camino de Valdeoliva, s/n
San Agustin del Guadalix
Madrid
28750
Spain

24 May 2024

Notified Body Confirmation Letter
Reference: EU2023-607/ 764888 rev 1

To whom it may concern,

Confirmation of the status of a formal application, written agreement, and appropriate surveillance in the framework of Regulation (EU) 2023/607 amending Regulations (EU) 2017/745 and (EU) 2017/746 as regards the transitional provisions for certain medical devices and in vitro diagnostic medical devices

This letter confirms that, **BSI Group The Netherlands B.V.**, a Notified Body (NB) designated against Regulation (EU) 2017/745 (MDR) and identified by the number **2797** on NANDO, has received a formal application in accordance with Section 4.3, first subparagraph of Annex VII of MDR and has signed a written agreement in accordance with Section 4.3, second subparagraph of Annex VII of MDR with the following manufacturer:

Becton Dickinson S.A.
Camino de Valdeoliva, s/n
San Agustin del Guadalix
Madrid
28750
Spain
SRN Number: ES-MF-000016479

BSI Group The Netherlands B.V.
Say Building
John M. Keynesplein 9, 1066 EP
Amsterdam, The Netherlands

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Validity of this letter may be verified by writing to Certificate.Verification@bsigroup.com

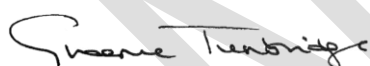
The devices covered by the formal application and the written agreement mentioned above are identified in the Tables below. Table 1 identifies the devices for which an MDR application has been received, written agreement concluded and for which the NB is also responsible for appropriate surveillance of the corresponding devices under the applicable Directive. Table 2 identifies the devices for which an MDR application has been received and a written agreement concluded, but the NB has not yet taken the responsibility for appropriate surveillance of the corresponding devices under the applicable Directive.

In the case of devices covered by certificates issued under Directive 90/385/EEC (AIMDD) or Directive 93/42/EEC (MDD) that expired after 26 May 2021 and before 20 March 2023, without having been withdrawn, this letter also confirms that the manufacturer signed the written agreement under MDR by the date of MDD/AIMDD certificate expiry; or provided evidence that a competent authority of a Member State had granted a derogation or exemption from the applicable conformity assessment procedure in accordance with Article 59(1) of MDR or Article 97(1) of the MDR respectively, by the 20 Mar 2023 for the relevant devices.

The transition timelines that apply to the devices covered by this letter, subject to the manufacturer's continued compliance to the other conditions specified in Article 120.3c of MDR (as amended by (EU) 2023/607), are shown below:

- 26 May 2026 for Class III custom-made implantable devices
- 31 December 2027 for Class III devices and Class IIb implantable devices excluding Well-established technologies (WET - sutures, staples, dental fillings, dental braces, tooth crowns, screws, wedges, plates, wires, pins, clips and connectors)
- 31 December 2028 for other Class IIb devices, Class IIa, Class I devices placed on the market in sterile condition or have a measuring function
- 31 December 2028 for devices not requiring the involvement of a notified body under MDD but requiring it under MDR (e.g., class I devices that qualify as re-usable surgical instruments)

On behalf of BSI Group The Netherlands B.V.,



Graeme Tunbridge
Senior Vice President, Medical Devices

Table 1: Devices covered by this letter and for which the NB is also responsible for appropriate surveillance of the corresponding devices under the applicable Directive:

Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified at the pre-application stage)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
N/A	N/A	N/A	N/A

Table 2: Devices covered by this letter and for which the NB is NOT responsible for appropriate surveillance of the corresponding devices under the applicable Directive:

Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified at the pre-application stage)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
BD Plastipak™ Luer-Lok™ Syringe (without needle) Basic UDI-DI: 038290SYLPEAGDM5 Listed on MDD certificate as: <i>Three-piece syringes without needle;</i> <i>Sterile BD Plastipak™ Luer-Lok™ syringes;</i> <i>Amber color syringes of: 50 ml</i>	Class IIa	N/A	Certificate reference: 95 06 0005 CP; expiry date 2024-05-24, Number: 0318
BD Plastipak™ Luer-Lok™ Convenience Pack (without needle) Basic UDI-DI: 038290UOKRIXNENG Listed on MDD certificate as: <i>Three-piece syringes without needle;</i> <i>Sterile BD Plastipak™ Luer-Lok™ syringes;</i>	Class IIa	'N/A'	Certificate reference: 95 06 0005 CP; expiry date 2024-05-24, NB number 0318

Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified at the pre-application stage)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
<i>Sizes:</i> <i>Convenience Pack of syringes of: 20 ml,</i> <i>Convenience Pack of syringes of: 50 ml</i>			
BD Plastipak™ Catheter Tip Syringe Basic UDI-DI: 038290TWOWNMYHW3 Listed on the MDD certificate as: <i>Syringes without needle, Insulin syringes; Three-piece sterile BD Plastipak™ syringes without needle; Syringes with Catheter Tip connection</i> <i>Sizes:</i> <i>Syringes of: 50 ml</i> <i>Syringes of: 100 ml</i>	Class I device placed on the market in sterile condition Class I device with a measuring function	'N/A'	Certificate reference: 2000 06 0273 CP; expiry date 2024-05-24, NB number 0318
BD Plastipak™ Catheter Tip Syringe (Bulk Non Sterile) Basic UDI-DI: 038290IHOCSHLU8L Listed on the MDD certificate as: <i>Syringes without needle; Non-sterile three-piece syringes without needle BD Plastipak™; Syringes with Catheter Tip connection</i> <i>Size:</i> <i>Syringes of: 50 ml</i>	Class I device with a measuring function	'N/A'	Certificate reference: 2019 09 0898 CP; expiry date 2024-05-24, NB# 0318
BD Plastipak™ Luer-Slip Syringe (without needle)	Class I device placed on the	'N/A'	Certificate reference:

Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified at the pre- application stage)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
Basic UDI-DI: 038290SRGVPAAEKC Listed on the MDD certificate as: <i>Syringes without needle, Insulin syringes; Three-piece sterile BD Plastipak™ syringes without needle; Luer-Slip syringes</i> <i>Sizes:</i> <i>Syringes of: 1 ml</i> <i>Syringes of: 2 ml</i> <i>Syringes of: 5 ml</i> <i>Syringes of: 10 ml</i> <i>Syringes of: 20 ml</i> <i>Syringes of: 30 ml</i> <i>Syringes of: 50 ml</i>	market in sterile condition Class I device with a measuring function		2000 06 0273 CP; expiry date 2024-05-24, NB number 0318
BD Plastipak™ Luer-Slip Syringe (without needle Bulk Non Sterile) Basic UDI-DI: 038290XOKHQIDL Listed on the MDD certificate as: <i>Syringes without needle; Non-sterile three-piece syringes without needle BD Plastipak™; Luer-Slip syringes</i> <i>Sizes:</i> <i>Syringes of: 1 ml</i> <i>Syringes of: 20 ml</i> <i>Syringes of: 50 ml</i>	Class I device with a measuring function	'N/A'	Certificate reference: 2019 09 0898 CP; expiry date 2024-05-24, NB# 0318
BD Whitacre Spinal Needle	Class III	'N/A'	Certificate reference:

Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified at the pre- application stage)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
Basic UDI-DI: 038290JPCVIFOCBR Listed on the MDD certificate as: <i>Spinal needles; Whitacre Pencil Point Spinal Needles; Sterile Whitacre Pencil Point Spinal Needle: BD Whitacre Needle</i> <i>Sizes:</i> 22G × 3.50" (0.7 × 90 mm) 24G × 3.50" (0.55 × 90 mm) 25G × 3.50" (0.50 × 90 mm) 27G × 3.50" (0.40 × 90 mm)			95 06 0005 CP and 2010 02 0700 ET; expiry date 2024-05-24, NB# 0318
BD Whitacre Spinal Needle Set (Spinal needle + introducer) Basic UDI-DI: 038290KITVJVYYLK Listed on the MDD certificates as: <i>Sterile Whitacre Pencil Point Spinal Needle with introducer set, BD Whitacre Needle</i> <i>Sizes:</i> 25G × 3.50" (0.50 × 90 mm) with introducer 20G × 1.25" (0.9 × 32 mm) 25G × 4.06" (0.50 × 103 mm) with introducer 20G × 1.25" (0.9 × 32 mm) 27G × 3.50" (0.40 × 90 mm) with introducer 22G × 1.25" (0.7 × 32 mm) 27G × 4.06" (0.40 × 103 mm) with introducer 22G × 1.25" (0.7 × 32 mm)	Class III	'N/A'	Certificate reference: 95 06 0005 CP and 2010 02 0700 ET; expiry date 2024-05-24, NB# 0318, to be read in conjunction with amending AEMPS letter dated 16 November 2021, ON0318/GH/JC/PS/95 04 0005
BD Quincke Spinal Needle	Class III	'N/A'	Certificate reference:

Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified at the pre- application stage)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
Basic UDI-DI: 038290HFPYXWDAGC Listed on the MDD certificate as: <i>Spinal needles; Spinal needle Quincke type point; Sterile spinal needle Quincke type point: BD Spinal Needle</i> <i>Sizes:</i> 18G × 3.00" (1.2 × 75 mm) 18G × 3.50" (1.2 × 90 mm) 19G × 3.00" (1.1 × 75 mm) 19G × 3.50" (1.1 × 90 mm) 20G × 1.50" (0.90 × 38 mm) 20G × 3.00" (0.90 × 75 mm) 20G × 3.50" (0.90 × 90 mm) 22G × 1.50" (0.7 × 38 mm) 22G × 2.50" (0.7 × 63 mm) 22G × 3.00" (0.7 × 75 mm) 22G × 3.50" (0.7 × 90 mm) 23G × 3.50" (0.64 × 90 mm) 25G × 3.00" (0.50 × 75 mm) 25G × 3.50" (0.50 × 90 mm) 26G × 3.50" (0.45 × 90 mm) 27G × 3.50" (0.40 × 90 mm)			95 06 0005 CP and 2010 02 0701 ET; expiry date 2024- 05-24, NB# 0318
BD Quincke Spinal Needle Set (Spinal needle + introducer) Basic UDI-DI: 038290JBDWOAQF5Y Listed on the MDD certificate as:	Class III	'N/A'	Certificate reference: 95 06 0005 CP and 2010 02 0701 ET; expiry date 2024- 05-24, NB# 0318, to be read in conjunction with amending AEMPS letter dated 15

Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified at the pre- application stage)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
<i>Spinal needles; Spinal needle Quincke type point; Sterile spinal needle Quincke type point with introducer set: BD Spinal Needle</i> <i>Sizes:</i> <i>25G × 3.50" (0.50 × 90 mm) con introducer / with introducer 20G × 1.25" (0.9 × 32 mm)</i> <i>26G × 3.50" (0.45 × 90 mm) con introducer / with introducer 20G × 1.25" (0.9 × 32 mm)</i> <i>27G × 3.50" (0.40 × 90 mm) con introducer / with introducer 22G × 1.25" (0.7 × 32 mm)</i>			November 2021 S/REF CNCps ID: 90811
BD Quincke Spinal NREFit™ Needle Basic UDI-DI: 038290WAHENDUTA4 Listed on the MDD certificate as: <i>Spinal needles; BD Quincke Spinal NREFit™ Needles; Sterile BD Quincke Spinal NREFit™ Needles BD Quincke Spinal NREFit™ Needle</i> <i>Sizes:</i> <i>25G × 3.00" (0.5 × 76.2 mm)</i> <i>18G × 3.00" (1.2 × 76.2 mm)</i> <i>18G × 3.50" (1.2 × 88.9 mm)</i> <i>22G × 1.50" (0.7 × 38.1 mm)</i> <i>22G × 2.50" (0.7 × 63.5 mm)</i> <i>22G × 3.00" (0.7 × 76.2 mm)</i> <i>22G × 3.50" (0.7 × 88.9 mm)</i> <i>25G × 3.50" (0.5 × 88.9 mm)</i> <i>25G × 2.00" (0.5 × 50.8 mm)</i> <i>19G × 3.00" (1.1 × 76.2 mm)</i>	Class III	N/A	Certificate reference: 95 06 0005 CP and 2020 04 0913 ET; expiry date 2024- 05-24, NB# 0318

Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified at the pre-application stage)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
19G × 3.50" (1.1 × 88.9 mm) 20G × 1.50" (0.9 × 38.1 mm) 20G × 3.00" (0.9 × 76.2 mm) 20G × 3.50" (0.9 × 88.9 mm) 23G × 3.50" (0.6 × 88.9 mm) 26G × 3.50" (0.45 × 88.9 mm) 27G × 3.50" (0.4 × 88.9 mm) 18G × 6.00" (1.20 × 152.4 mm) 20G × 6.00" (0.90 × 152.4 mm) 22G × 5.00" (0.70 × 127 mm) 22G × 7.00" (0.70 × 177.8 mm) 25G × 4.70" (0.50 × 119.1 mm)			
BD Quincke Spinal NRFit™ Needle Set (NRFit Spinal Needle + introducer) Basic UDI-DI: 038290MIVCNTRJEZ Listed on the MDD certificate as: <i>Spinal needles; BD Quincke Spinal NRFit™ Needles; Sterile BD Quincke spinal NRFit™ Needle with introducer set. BD Quincke Spinal NRFit™ Needle Set</i> <i>Sizes:</i> <i>25G × 3.50" (0.50 × 88.9 mm) con introducer / with introducer 20 G × 1.25" (0.90 × 31.8 mm)</i> <i>27G × 3.50" (0.40 × 88.9 mm) con introducer / with introducer 22 G × 1.25" (0.70 × 31.8 mm)</i>	Class III	N/A	Certificate reference: 95 06 0005 CP and 2020 04 0913 ET; expiry date 2024-05-24, NB# 0318, to be read in conjunction with amending AEMPS letter dated 15 november 2021, S/REF CNCps ID: 92262

Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified at the pre- application stage)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
<i>26G × 3.50" (0.45 × 88.9 mm) with introducer / with introducer 20G × 1.25" (0.90 × 31.8 mm)</i>			
BD Whitacre Spinal NRFit™ Needle Basic UDI-DI: 038290RFMMCXCCX Listed on the MDD certificate as: <i>Spinal needles; BD Whitacre Spinal NRFit™ Needles; Sterile BD Whitacre Spinal NRFit™ Needle BD Whitacre Spinal NRFit™ Needle</i> <i>Sizes:</i> <i>25G × 4.06" (0.50 × 103.2 mm)</i> <i>25G × 4.70" (0.50 × 119.1 mm)</i> <i>27G × 4.06" (0.40 × 103.2 mm)</i> <i>27G × 4.70" (0.40 × 119.1 mm)</i> <i>22G × 3.50" (0.70 × 88.9 mm)</i> <i>24G × 3.50" (0.55 × 88.9 mm)</i> <i>25G × 3.50" (0.50 × 88.9 mm)</i> <i>27G × 3.50" (0.40 × 88.9 mm)</i>	Class III	N/A	Certificate reference: 95 06 005 CP and 2020 04 0912 ET; expiry date 2024- 05-24, NB# 0318
BD Whitacre Spinal NRFit™ Needle Set (NRFit Spinal Needle + introducer) Basic UDI-DI: 038290EDURMMPSBE Listed on the MDD certificate as: <i>Spinal needles; BD Whitacre Spinal NRFit™ Needles; Sterile BD Whitacre Spinal NRFit™ Needle with introducer</i>	Class III	N/A	Certificate reference: 95 06 005 CP and 2020 04 0912 ET; expiry date 2024- 05-24, NB# 0318

Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified at the pre-application stage)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
<i>set. BD Whitacre Spinal NRFit™ Needle Set.</i> <i>Sizes:</i> <i>27G × 3.50" (0.40 × 88.9 mm) with introducer 22G × 1.25" (0.70 × 31.8 mm)</i> <i>25G × 3.50" (0.50 × 88.9 mm) with introducer 20G × 1.25" (0.90 × 31.8 mm)</i> <i>25G × 4.06" (0.50 × 103.2 mm) with introducer 20G × 1.25" (0.90 × 31.8 mm)</i> <i>27G × 4.06" (0.40 × 103.2 mm) with introducer 22G × 1.25" (0.70 × 31.8 mm)</i>			
BD Spinal Introducer NRFit Needle Basic UDI-DI (Sterile): 038290XGKVYVIXRA Listed on the MDD certificate as: <i>Needle introducer; Sterile BD Spinal needle introducer NRFit™ BD Spinal needle introducer NRFit™</i> <i>Sizes:</i> <i>20G × 1.25" (0.90 × 31.8 mm)</i> <i>22G × 1.25" (0.70 × 31.8 mm)</i>	Class IIa	N/A	Certificate reference: 95 06 0005 CP; expiry date 2024-05-24, NB# 0318
BD Discardit II Syringe (with Needle) Basic UDI-DI (Microlance needle): 038290HLQCKJFL8D Listed on the MDD certificate as:	Class IIa	N/A	Certificate reference: 95 06 0006 CP; expiry date 2024-05-26, NB# 0318 MDD Certificate issued to BECTON DICKINSON S.A.

Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified at the pre- application stage)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
<i>Syringes with needle; Sterile two-piece syringes with needle BD Discardit™ II</i> <i>Sizes:</i> <i>Syringes of: 2 ml:</i> <i>With needle: 21G × 5/8" (0.8 × 16 mm)</i> <i>With needle: 21G × 1 1/2" (0.8 × 40 mm)</i> <i>With needle: 22G × 1 1/2" (0.7 × 40 mm)</i> <i>With needle: 23G × 1" (0.6 × 25 mm)</i> <i>With needle: 24G × 1" (0.55 × 25 mm)</i> <i>Syringes of: 5 ml</i> <i>With needle: 21G × 1 1/2" (0.8 × 40 mm)</i> <i>With Needle: 23G 1" (0.6 × 25 mm)</i> <i>Syringes of: 10 ml</i> <i>With needle: 21G × 1 1/2" (0.8 × 40 mm)</i> <i>With needle: 21G × 1" (0.8 × 25 mm)</i> <i>Syringes of: 20 ml</i> <i>With needle: 21G × 1 1/2" (0.8 × 40 mm)</i>			C/ Mequinenza, s/n. E-22520 FRAGA (Huesca) España (Part of the same larger organisation)
BD Discardit II Non Sterile Bulk Syringe (without needle) Basic UDI-DI: 038290IWCNEZKAT Listed on the MDD certificate as:	Class I device with a measuring function	N/A	Certificate reference: 99 03 0213 CP; expiry date 2024-05-26, NB# 0318 MDD Certificate issued to BECTON DICKINSON S.A.

Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified at the pre-application stage)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
<i>Syringes without needle; Non-sterile syringes without needle; Two-piece syringes BD Discardit™ II</i> sizes: <i>Syringes of: 2 ml</i> <i>Syringes of: 5 ml</i> <i>Syringes of: 10 ml</i> <i>Syringes of: 20 ml</i>			C/ Mequinenza, s/n. E-22520 FRAGA (Huesca) España (Part of the same larger organisation)
BD Discardit II Syringe (without needle) Basic UDI-DI: 038290DDEODFOLXL Listed on the MDD certificate as: <i>Syringes without needle and syringes with blunt fill needle; Sterile syringes without needle; Two-piece syringes BD Discardit™ II</i> <i>Syringes of: 2 ml</i> <i>Syringes of: 5 ml</i> <i>Syringes of: 10 ml</i> <i>Syringes of: 20 ml</i>	Class I device placed on the market in sterile condition Class I device with a measuring function	N/A	Certificate reference: 2000 06 0272 CP; expiry date 2024-05-26, NB# 0318 MDD Certificate issued to BECTON DICKINSON S.A. C/ Mequinenza, s/n. E-22520 FRAGA (Huesca) España (Part of the same larger organisation)
BD Microlance™ 3 Needle Basic UDI-DI: 038290MLFUVFGJET Listed on the MDD certificate as: <i>Needles; Sterile hypodermic needles BD Microlance™ 3</i> Sizes: <i>Needle 20G × 1 ½" (0.9 × 40 mm)</i> <i>Needle 20G × 1" (0.9 × 25 mm)</i> <i>Needle 21G × 1" (0.8 × 25 mm)</i>	Class IIa	N/A	Certificate reference: 95 06 0006 CP; expiry date 2024-05-26, NB# 0318 MDD Certificate issued to BECTON DICKINSON S.A. C/ Mequinenza, s/n. E-22520 FRAGA (Huesca) España (Part of the same larger organisation)

Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified at the pre- application stage)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
<i>Needle 21G × 1 ½" (0.8 × 40 mm)</i> <i>Needle 21G × ⅝" (0.8 × 16 mm)</i> <i>Needle 22G × 1 ¼" (0.7 × 30 mm)</i> <i>Needle 22G × 1 ½" (0.7 × 40 mm)</i> <i>Needle 22G × 1" (0.7 × 25 mm)</i> <i>Needle 23G × 1 ¼" (0.6 × 30 mm)</i> <i>Needle 23G × 1" (0.6 × 25 mm)</i> <i>Needle 22G × 2" (0.7 × 50 mm)</i> <i>Needle 26G × ⅜" (0.45 × 10 mm)</i> <i>Needle 25G × 1" (0.5 × 25 mm)</i> <i>Needle 25G × ⅝" (0.5 × 16 mm)</i> <i>Needle 27G × ½" (0.4 × 13 mm)</i> <i>Needle 16G × 1 ½" (1.6 × 40 mm)</i> <i>Needle 21G × 2" (0.8 × 50 mm)</i> <i>Needle 19G × 1 ½" (1.1 × 40 mm)</i> <i>Needle 19G × 1" (1.1 × 25 mm)</i> <i>Needle 19G × 2" (1.1 × 50 mm)</i> <i>Needle 18G × 2" (1.2 × 50 mm)</i> <i>Needle 27G × ¾" (0.4 × 19 mm)</i> <i>Needle 26G × ½" (0.45 × 13 mm)</i> <i>Needle 30G × ½" (0.3 × 13 mm)</i> <i>Needle 24G × 1" (0.55 × 25 mm)</i> <i>Needle 26G × ⅝" (0.45 × 16 mm)</i> <i>Needle 18G × 1 ½" (1.2 × 40 mm)</i> <i>Needle 27G × ⅜" (0.4 × 10 mm)</i> <i>Needle 23G × 1 ½" (0.6 × 40 mm)</i>			
BD Microlance TM 3 Non Sterile Bulk Needle	Class IIa	N/A	Certificate reference: 95 06 0006 CP; expiry date 2024-05-26, NB# 0318, to be

Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified at the pre- application stage)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
Basic UDI-DI: 038290MVVIHDPPLC Listed on the MDD certificate as: <i>Needles; Non-sterile hypodermic needles BD Microlance™ 3</i> <i>Needle 25G × 5/8" (0.5 × 16 mm)</i> <i>Needle 21G × 5/8" (0.8 × 16 mm)</i> <i>Needle 25G × 1" (0.5 × 25 mm)</i> <i>Needle 21G × 1" (0.8 × 25 mm)</i> <i>Needle 21G × 1 1/2" (0.8 × 40 mm)</i> <i>Needle 20G × 1 1/2" (0.9 × 40 mm)</i> <i>Needle 22G × 1 1/4" (0.7 × 30 mm)</i> <i>Needle 30G × 1/2" (0.3 × 13 mm)</i> <i>Needle 27G × 1/2" (0.4 × 13 mm)</i> <i>Needle 19G × 1 1/2" (1.1 × 40 mm)</i> <i>Needle 14G × 1 1/4" (2.1 × 30 mm)</i> <i>Needle 26G × 1/2" (0.45 × 13 mm)</i> <i>Needle 26G × 3/8" (0.45 × 10 mm)</i> <i>Needle 23G × 1" (0.6 × 25 mm)</i> <i>Needle 21G × 2" (0.8 × 50 mm)</i> <i>Needle 18G × 2" (1.2 × 50 mm)</i> <i>Needle 27G × 3/4" (0.4 × 19 mm)</i> <i>Needle 24G × 1" (0.55 × 25 mm)</i> <i>Needle 18G × 1 1/2" (1.2 × 40 mm)</i> <i>Needle 22G × 1" (0.7 × 25 mm)</i> <i>Needle 22G × 1 1/2" (0.7 × 40 mm)</i> <i>Needle 23G × 1 1/4" (0.6 × 30 mm)</i> <i>New non-sterile variant of the Microlance™ 3 27G × 3/8" (0.4 × 10 mm) needle</i>			read in conjunction with amending AEMPS letter dated 10 March 2022 regarding and change submitted 22 December 2021, S/REF 94331 MDD Certificate issued to BECTON DICKINSON S.A. C/ Mequinenza, s/n. E-22520 FRAGA (Huesca) España (Part of the same larger organisation)

Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified at the pre- application stage)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
BD SoloShot™ Mini Syringe Basic UDI-DI: 038290CDXR0BXD6 Listed on the MDD certificate as: <i>Syringes with needle; Sterile auto-disable two-piece syringes with integrated cannula BD SoloShot™ Mini</i> <i>Sizes:</i> <i>Syringes of: 0.5 ml</i> <i>With needle: 23G × 1" (0.6 × 25 mm)</i> <i>With needle: 24G × ¾" (0.55 × 19 mm)</i> <i>With needle: 25G × ⅝" (0.5 × 16 mm)</i> <i>With needle: 25G × 1" (0.5 × 25 mm)</i> <i>Syringes of: 0.1 ml</i> <i>With needle: 27G × ⅜" (0.4 × 10 mm)</i> <i>Syringes of: 0.05 ml</i> <i>With needle: 27G × ⅜" (0.4 × 10 mm)</i> <i>Syringes of 0.2 ml</i> <i>With needle: 23G × 1" (0.6 × 25 mm)</i> <i>Syringes of 0.3 ml</i> <i>With needle: 23G × 1" (0.6 × 25 mm)</i> <i>Syringes of 0.4 ml</i> <i>With needle: 23G × 1" (0.6 × 25 mm)</i>	Class IIa	N/A	Certificate reference: 95 06 0006 CP; expiry date 2024-05-26, NB# 0318 MDD Certificate issued to BECTON DICKINSON S.A. C/ Mequinenza, s/n. E-22520 FRAGA (Huesca) España (Part of the same larger organisation)

Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified at the pre- application stage)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
BD SoloShot™ IX Syringe Basic UDI-DI: 038290MJTMIQJVL Listed on the MDD certificate as: <i>Syringes with needle; Sterile auto-disable two piece syringes with integrated cannula BD SoloShot™ IX</i> <i>Sizes:</i> <i>Syringes of: 0.5 ml</i> <i>With needle: 23G × 1" (0.6 × 25 mm)</i> <i>With needle: 24G × ¾" (0.55 × 19 mm)</i> <i>With needle: 24G × 1" (0.55 × 25 mm)</i> <i>With needle: 25G × ⅝" (0.5 × 16 mm)</i> <i>With needle: 25G × 1" (0.5 × 25 mm)</i> <i>Syringes of: 1 ml</i> <i>With needles: 22G × 1" (0.7 × 25 mm)</i> <i>With needle: 23G × 1" (0.6 × 25 mm)</i>	Class IIa	N/A	Certificate reference: 95 06 0006 CP; expiry date 2024-05-26, NB# 0318 MDD Certificate issued to BECTON DICKINSON S.A. C/ Mequinenza, s/n. E-22520 FRAGA (Huesca) España (Part of the same larger organisation)
BD Flu+ Syringe Basic UDI-DI: 038290VFCSOUDWGE Listed on the MDD certificate as: <i>Syringes with needle; Sterile two-piece syringes with integrated cannula BD Flu+</i> <i>Sizes:</i>	Class IIa	N/A	Certificate reference: 95 06 0006 CP; expiry date 2024-05-26, NB# 0318, to be read in conjunction with amending AEMPS letter dated 20 October 2021 regarding and change submitted 20 August 2021, S / REF: 90802 MDD Certificate issued to BECTON DICKINSON S.A., C/ Mequinenza, s/n. E-22520

Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified at the pre-application stage)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
<i>BD® Flu+ Syringe 0.25 - 1 mL with needle: 23G × 1" (0.6 × 25 mm)</i> <i>BD® Flu+ Syringe 0.25 - 1 mL with needle: 25G × 5/8" (0.5 × 16 mm)</i> <i>BD® Flu+ Syringe 0.25 - 1 mL with needle: 25G × 1" (0.5 × 25 mm)</i> <i>BD® Flu+ Syringe 0.1 - 1 mL with needle: 23G × 1" (0.6 × 25 mm) 0.1 – 1 ml</i> <i>BD® Flu+ Syringe 0.1 - 1 mL with needle: 25G × 1" (0.5 × 25 mm) 0.1 – 1 ml</i>			FRAGA (Huesca) España (Part of the same larger organisation)
BD Emerald TM Syringe (without needle) Basic UDI-DI: 038290PDVJVUZUKV Listed on the MDD certificate as: <i>Syringes without needle and syringes with blunt fill needle; Sterile syringes without needle; Sterile three-piece syringes BD Emerald™</i> <i>Sizes:</i> <i>Syringes of: 2 ml</i> <i>Syringes of: 5 ml</i> <i>Syringes of: 10 ml</i>	Class I device placed on the market in sterile condition Class I device with a measuring function	N/A	Certificate reference: 2000 06 0272 CP; expiry date 2024-05-26, NB# 0318 MDD Certificate issued to BECTON DICKINSON S.A., C/ Mequinenza, s/n. E-22520 FRAGA (Huesca) España (Part of the same larger organisation)
BD Emerald™ Syringe (with needle) Basic UDI-DI (Microlance needle): 038290JGXPPFFCE5 Listed on the MDD certificate as: <i>Sterile three-piece syringes with needle BD Emerald™;</i>	Class IIa	N/A	Certificate reference: 95 06 0006 CP; expiry date 2024-05-26, NB# 0318 MDD Certificate issued to BECTON DICKINSON S.A., C/ Mequinenza, s/n. E-22520 FRAGA (Huesca) España (Part of the same larger organisation)

Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified at the pre- application stage)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
<i>Syringes of 2 ml and needle side by side:</i> <i>With needle: 21G × 1 ½" (0.8 × 40 mm)</i> <i>With needle: 23G × 1" (0.6 × 25 mm)</i> <i>Syringes of 2 ml and pre-attached needle</i> <i>With needle: 21G × 1 ½" (0.8 × 40 mm)</i> <i>With needle: 23G × 1" (0.6 × 25 mm)</i> <i>With needle: 25G × ⅝" (0.5 × 16 mm)</i> <i>Syringes of 5 ml and needle side by side</i> <i>With needle: 21G × 1 ½" (0.8 × 40 mm)</i> <i>With needle: 23G × 1" (0.6 × 25 mm)</i> <i>With needle: 24G × 1" (0.55 × 25 mm)</i> <i>Syringes of 5 ml and pre-attached needle</i> <i>With needle: 21G × 1 ½" (0.8 × 40 mm)</i> <i>With needle: 23G × 1" (0.6 × 25 mm)</i> <i>Syringes of 10 ml and needle side by side</i> <i>With needle: 21G × 1 ½" (0.8 × 40 mm)</i> <i>With needle: 21G × 1" (0.8 × 25 mm)</i>			

Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified at the pre-application stage)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
<i>Syringes of 10 ml and pre-attached needle</i> <i>With needle: 21G × 1 ½" (0.8 × 40 mm)</i>			
BD Emerald™ Non Sterile Bulk Syringe (without needle) Basic UDI-DI: 038290ZDNJQVHDJV Listed on the MDD certificate as: <i>Syringes without needle; Non-sterile syringes without needle; Three-piece bulk syringes BD Emerald™</i> <i>Sizes:</i> <i>Syringes of: 2 ml</i> <i>Syringes of: 5 ml</i> <i>Syringes of: 10 ml</i>	Class I device with a measuring function	N/A	Certificate reference: 99 03 0213 CP; expiry date 2024-05-26, NB# 0318 MDD Certificate issued to BECTON DICKINSON S.A., C/ Mequinenza, s/n. E-22520 FRAGA (Huesca) España (Part of the same larger organisation)
BD Emerald™ PRO Syringe (with needle) Basic UDI-DI (Microlance needle): 038290GRPEWKVVF5 Listed on the MDD certificate as: <i>Sterile three-piece syringes with reuse prevention and needle BD Emerald™ PRO;</i> <i>Syringes of 2 ml and needle side by side</i> <i>With needle: 21G × 1 ½" (0.8 × 40 mm)</i> <i>With needle: 23G × 1" (0.6 × 25 mm)</i>	Class IIa	N/A	Certificate reference: 95 06 0006 CP; expiry date 2024-05-26, NB# 0318 MDD Certificate issued to BECTON DICKINSON S.A., C/ Mequinenza, s/n. E-22520 FRAGA (Huesca) España (Part of the same larger organisation)

Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified at the pre-application stage)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
<i>Syringes of 5 ml and needle side by side</i> <i>With needle: 21G × 1 ½" (0.8 × 40 mm)</i> <i>With needle: 23G × 1" (0.6 × 25 mm)</i> <i>Syringes of 10 ml and needle side by side</i> <i>With needle: 21G × 1 ½" (0.8 × 40 mm)</i>			
BD Blunt Fill Needle Basic UDI-DI: 38290NGGTJRVACW Listed on the MDD certificate as: <i>Blunt fill needle; Sterile needle to aspirate pharmaceutical fluids from vials or ampoules; Needle BD Blunt Fill Needle</i> <i>Size: 18G × 1 ½" (1.2 × 40 mm)</i>	Class I device placed on the market in sterile condition	N/A	Certificate reference: 2015 03 0838 CP; expiry date 2024-05-26, NB# 0318 MDD Certificate issued to BECTON DICKINSON S.A., C/ Mequinenza, s/n. E-22520 FRAGA (Huesca) España (Part of the same larger organisation)
BD Plastipak Insulin syringe (with Microlance Needle) Basic UDI: 038290HVAFTJLHAH Listed on the MDD certificate as: <i>Three-piece insulin Luer-Slip syringes with needle;</i> <i>Sizes:</i>	Class IIa	N/A	Certificate reference: 95 06 0005 CP; expiry date 2024-05-24, Number: 0318

Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified at the pre-application stage)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
<i>Sterile BD Plastipak™ Insulin 1ml 40 I.U. syringes With needle: 30G × ½" (0.3 × 13 mm)</i> <i>Sterile BD Plastipak™ Insulin 1ml 100 I.U. syringes</i> <i>With needle: 25G × ⅝" (0.5 × 16 mm)</i> <i>With needle: 26G × ⅝" (0.45 × 10mm)</i>			
BD Plastipak Insulin syringe (without needle) Basic UDI: 038290HUGAZKNPCU Listed on the MDD certificate as: <i>Three-piece sterile BD Plastipak™ syringes without needle; Luer-Slip insulin syringes;</i> <i>Size:</i> <i>Insulin 1ml 40 I.U. syringes, Insulin 1ml 100 I.U. syringes</i>	Class I device placed on the market in sterile condition Class I device with a measuring function	N/A	Certificate reference: 2000 06 0273 CP; expiry date 2024-05-24, NB number 0318
BD Discardit II Syringe with PrecisionGlide Needle Basic UDI: 038290CTTJUWJFHH Listed on the MDD certificate as: <i>Sterile Hypodermic Syringes for Single use with needle (2 piece)</i> <i>Sizes: 2ml, 5ml</i>	Class IIa	N/A	Certificate reference: 11317-2017-CE-IND-NA-PS; expiry date 2024-05-27, NB number 2460
BD Emerald™ Syringe	Class IIa	N/A	Certificate reference:

Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified at the pre- application stage)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
(with needle) Basic UDI-DI (Microlance needle): 038290JGXPPFFCE5 Listed on the MDD certificate as: <i>Sterile Hypodermic Syringes for Single use with needle (2 piece)</i> <i>Sizes: 2ml, 5ml, 10ml,</i>			11317-2017-CE-IND-NA-PS; expiry date 2024-05-27, NB number 2460
BD Emerald™ PRO Syringe (with PrecisionGlide needle) Basic UDI: 038290CRXJAJPPCU Listed on the MDD certificate as: <i>Sterile Hypodermic Syringes for Single use with needle (2 piece)</i> <i>Sizes: 2ml, 3ml, 5ml, 10ml</i>	Class IIa	N/A	Certificate reference: 11317-2017-CE-IND-NA-PS; expiry date 2024-05-27, NB number 2460
BD Emerald™ Syringe (with PrecisionGlide Needle) Basic UDI: 038290EIGAHXZP3W Listed on the MDD certificate as: <i>Sterile Hypodermic Syringes for Single use with needle (2 piece)</i> <i>Sizes: 2ml, 3ml, 5ml, 10ml</i>	Class IIa	N/A	Certificate reference: 11317-2017-CE-IND-NA-PS; expiry date 2024-05-27, NB# 2460

Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified at the pre- application stage)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
BD Quincke Spinal Needle Basic UDI: 038290BKWHSNSEBL Listed on the MDD certificate as: <i>BD Spinal needle Quincke type point;</i> <i>BD Spinal needle Quincke type point</i> 18 GA 6 IN sterile 20 GA 1.50 IN sterile 20 GA 6.00 IN sterile 22 GA 5.00 IN sterile 22 GA 7.00 IN sterile 25 GA 2.00 IN sterile 25 GA 4.11/16 IN sterile	Class III	N/A	Certificate reference: 2010 02 0698 ET, expires 17-12-2023, NB # 0318 MDD Certificate issued to Becton, Dickinson and Company 1 Becton Dr, Franklin Lakes, New Jersey 07417. USA (Part of the same larger organisation)
BD Whitacre Spinal Needle Basic UDI: 038290AGCTSBAT29 Listed on the MDD certificate as: <i>Anesthesia needles; BD Spinal needle</i> <i>Whitacre type point:</i> 22GA 3.50 IN sterile 25GA 4.11/16" Sterile 25GA 3.50 IN TW Sterile 25GA 5.00 IN TW Sterile 27GA 4.11/16" Sterile 27GA 3.50 IN Sterile 27GA 5.00 IN TW Sterile	Class III	N/A	Certificate reference: 2010 02 0699 ET, expires 17-12-2023, NB # 0318 MDD Certificate issued to Becton, Dickinson and Company 1 Becton Dr, Franklin Lakes, New Jersey 07417. USA (Part of the same larger organisation)
BD® Neonatal Spinal Needle Quincke Basic UDI: 038290HFPYXWDAGC	Class III	N/A	Certificate reference:

Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified at the pre- application stage)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
Listed on the MDD certificate as: <i>Spinal needles; Spinal needle Quincke type point; Sterile spinal needle Quincke type point: BD Spinal Needle</i> <i>Sizes:</i> <i>25G × 1.00" (0.50 × 25 mm) Neonatal</i> <i>25G × 2.00" (0.50 × 51 mm) Neonatal</i>			95 06 0005 CP and 2010 02 0701 ET; expiry date 2024-05-24, NB# 0318
BD® NRFit Spinal Needle Quincke Bulk non sterile Basic UDI: 038290HDECPCXJYQ Listed on the MDD certificate as: <i>Non-sterile BD Quincke Spinal NRFit™ Needles. BD Quincke Spinal NRFit™ Needle:</i> <i>22G × 3.50" (0,70 × 88.9 mm)</i> <i>22G × 2.50" (0,70 × 63.5 mm)</i> <i>22G × 7.00" (0,70 × 177.8 mm)</i> <i>18G × 3.50" (1.2 × 88.9 mm)</i> <i>22G × 1.50" (0.7 × 38.1 mm)</i> <i>22G × 5.00" (0.70 × 127 mm)</i> <i>25G × 2.00" (0.5 × 50.8 mm)</i> <i>25G × 3.50" (0.5 × 88.9 mm)</i> <i>20G × 3.50" (0.9 × 88.9 mm)</i> <i>26G × 3.50" (0.45 × 88.9 mm)</i> <i>27G × 3.50" (0.4 × 88.9 mm)</i>	Class III	N/A	Certificate reference: 95 06 0005 CP and 2020 04 0913 ET; expiry date 2024-05-24, NB# 0318
BD® NRFit Spinal Needle Whitacre	Class III	N/A	Certificate reference:

Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified at the pre- application stage)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
Basic UDI: 038290HJEEJKUH48 Listed on the MDD certificate as: <i>Non-sterile BD Whitacre Spinal NRFit™ Needles</i> <i>BD Whitacre Spinal NRFit™ Needle:</i> 25G × 4.06" (0.50 × 103.2 mm) 27G × 4.06" (0.40 × 103.2 mm) 22G × 3.50" (0.70 × 88.9 mm) 24G × 3.50" (0.55 × 88.9 mm) 25G × 3.50" (0.50 × 88.9 mm) 25G × 4.70" (0.50 × 119.1 mm) 27G × 3.50" (0.40 × 88.9 mm) 27G × 4.70" (0.40 × 119.1 mm)			95 06 0005 CP and 2020 04 0913 ET; expiry date 2024- 05-24, NB# 0318
BD® Spinal Introducer NRFit™ Needle (Bulk non-sterile): Basic UDI: 038290HXSFAVSKHF Listed on the MDD certificate as: <i>Needle introducer; Non-sterile BD Spinal needle introducer NRFit™; BD Spinal needle introducer NRFit™</i> 20G × 1.25" (0.90 × 31.8 mm) 22G × 1.25" (0.70 × 31.8 mm)	Class IIa	N/A	Certificate reference: 95 06 0005 CP; expiry date 2024-05-24, NB# 0318
BD Plastipak™ Luer-Lok™ Syringe with Blunt Fill Needle Basic UDI: 038290CYRAILLJCC Listed on the MDD certificate as:	Class IIa	N/A	Certificate reference: 95 06 0005 CP; expiry date 2024-05-24, NB# 0318

Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified at the pre- application stage)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
<i>Three-piece syringes with needle; Sterile BD Plastipak™ Luer-Lok™ syringes with Blunt Fill Needle</i> <i>Sizes:</i> <i>Syringes of 50 ml with needle 18G × 1½" (1.2 × 40 mm)</i> <i>Amber color syringes of 50 ml with needle 18G × 1½" (1.2 × 40 mm)</i> <i>Syringes of 50 ml with Needle 18G × 1" (1.2 × 25 mm)</i> <i>Amber color syringes of 50 ml with Needle 18G × 1" (1.2 × 25 mm)</i>			
BD Plastipak™ Luer-Lok™ Syringe (without needle) Basic UDI-DI: 38290SRGVPAAEKC Listed on the MDD certificate as: <i>Three-piece syringes without needle</i> <i>Sterile BD Plastipak™ Luer-Lok™ syringes</i> <i>Syringes of: 10 ml</i> <i>Syringes of: 20 ml</i> <i>Syringes of: 30 ml</i> <i>Syringes of: 50 ml</i>	Class IIa	N/A	Certificate reference: 95 06 0005 CP; expiry date 2024-05-24, NB# 0318
BD Plastipak™ Luer-Lok™ Syringes (without needle Bulk Non Sterile) Basic UDI-DI: 38290XOKHQIDL Listed on the MDD certificate as:	Class IIa	N/A	Certificate reference: 95 06 0005 CP; expiry date 2024-05-24, NB# 0318

Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified at the pre- application stage)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
<i>Three-piece syringes without needle; Non-sterile BD Plastipak™ Luer-Lok™ syringes</i> <i>Syringes of: 20 ml</i> <i>Syringes of: 50 ml</i>			
BD Emerald™ Syringe with Blunt Fill Needle Basic UDI: 038290DUTBGSTYEB Listed on the MDD certificate as: <i>Syringes without needle and syringes with blunt fill needle; Sterile three-piece syringes with blunt fill needle to aspirate pharmaceutical fluids from vials or ampoules; Syringe with blunt fill needle</i> <i>Syringe of 2 ml with 18G × 1 ½" (1.2 × 40 mm) blunt fill needle</i> <i>Syringe of 5 ml with 18G × 1 ½" (1.2 × 40 mm) blunt fill needle</i> <i>Syringe of 10 ml with 18G × 1 ½" (1.2 × 40 mm) blunt fill needle</i>	Class I device placed on the market in sterile condition Class I device with a measuring function	N/A	Certificate reference: 2000 06 0272 CP; expiry date 2024-05-26, NB# 0318 MDD Certificate issued to BECTON DICKINSON S.A. C/ Mequinenza, s/n. E-22520 FRAGA (Huesca) España (Part of the same larger organisation)
BD Emerald™ Syringe (without needle) Basic UDI: 038290ECEAGPPW9 Listed on the MDD certificate as: <i>Sterile Hypodermic Syringes for Single use without needle (2 piece)</i> Sizes:	Class I device placed on the market in sterile condition Class I device with a measuring function	N/A	Certificate reference: MDD/AIMDD Certificate #11317-2017-CE-IND-NA-PS; expiry date 2024-05-27, NB# 2460 MDD Certificate issued to: Becton Dickinson India Pvt. Ltd., Plot No. 1, Sector 3, IMT, Bawal – 123 501, District Rewari, Haryana, India.

Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified at the pre-application stage)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
2ml, 3ml, 5ml, 10ml			(Part of the same larger organisation)
BD Discadit II Syringe (without needle) Basic UDI: 038290BUVPCNDNEW Listed on the MDD certificate as: <i>Sterile Hypodermic Syringes for Single use without needle (2 piece)</i> <i>Sizes:</i> 2ml, 5ml	Class I device placed on the market in sterile condition Class I device with a measuring function	N/A	Certificate reference: 11317-2017-CE-IND-NA-PS; expiry date 2024-05-27, NB# 2460 MDD Certificate issued to: Becton Dickinson India Pvt. Ltd., Plot No. 1, Sector 3, IMT, Bawal – 123 501, District Rewari, Haryana, India. (Part of the same larger organisation)
N/A	N/A	N/A	N/A

Confirmation Letter Revision History

Date	Action
2024/03/12	Initial issue
2024/05/24	Amended to add devices