



TECHNICAL DATA SHEET

BD Microlance™ 3 hypodermic needle Sterile, Single Use, Latex free

1. General Information

1.1 General

Intended use: BD Microlance™ 3 hypodermic needles are single-use medical devices intended for the hypodermic administration of pharmaceuticals.



BD Microlance™ 3 hypodermic needles are manufactured in different sizes, depending on the various exterior diameters and lengths of the cannulas. Each type of needle is recognized by the colour of the hub and by the identification system, both by the International System of Units (measurement in millimetres) and the American system (measurements in inches).

BD conventional needle design, materials and clinical application are based on well-established technologies and procedures. Sterile, single use disposable Microlance needles have been manufactured by BD and used successfully for over 50 years. Microlance needles are designed to operate with other devices. The needle hub has a female luer fitting which mates to male luer fittings, and is compatible with luer slip or luer lock syringes.



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Gauge and Diameter information

| GAUGE | DIAMETER (mm) |
|--------------|----------------------|
| 14 / 2,1 mm | 1,950-2,150 |
| 16 / 1,6 mm | 1,600-1,690 |
| 18 / 1,2 mm | 1,200-1,300 |
| 19 / 1,1 mm | 1,030-1,100 |
| 20 / 0,9 mm | 0,860-0,920 |
| 21 / 0,8 mm | 0,800-0,830 |
| 22 / 0,7 mm | 0,698-0,730 |
| 23 / 0,6 mm | 0,600-0,673 |
| 24 / 0,55 mm | 0,550-0,580 |
| 25 / 0,5 mm | 0,500-0,530 |
| 26 / 0,45 mm | 0,440-0,470 |
| 27 / 0,4 mm | 0,400-0,420 |
| 30 / 0.3 mm | 0.298-0.320 |

SPECIAL NEEDLES

| BD Reference | Description Gauge/Inches | Length | Wall | Color code | Box (units) | Case (units) |
|---------------------|---------------------------------|---------------|-------------|-------------------|--------------------|---------------------|
| 304000 | 30G x ½" | 13 mm | Regular | Yellow | 100 | 5.000 |
| 304434 | 21G x 5/8" | 16 mm | Thin | Green | 100 | 5.000 |
| 301700 | 19G x 1" | 25 mm | Thin | Ivory | 100 | 5.000 |
| 301750 | 19G x 2" | 50 mm | Thin | Ivory | 100 | 4.000 |
| 300637 | 16G x 1½" | 40 mm | Regular | White | 100 | 5.000 |

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REGULAR NEEDLES

| BD Reference | Description Gauge/Inches | Length | Wall | Color code | Box (units) | Case (units) |
|---------------------|---------------------------------|---------------|-------------|-------------------|--------------------|---------------------|
| 302200 | 27G x 3/4" | 19 mm | Regular | Grey | 100 | 5.000 |
| 300635 | 27G x 1/2" | 13 mm | Regular | Grey | 100 | 5.000 |
| 304300 | 26G x 5/8" | 16 mm | Regular | Brown | 100 | 5.000 |
| 303800 | 26G x 1/2" | 13 mm | Regular | Brown | 100 | 5.000 |
| 300300 | 26G x 3/8" | 10 mm | Regular | Brown | 100 | 5.000 |
| 300400 | 25G x 1" | 25 mm | Regular | Orange | 100 | 5.000 |
| 300600 | 25G x 5/8" | 16 mm | Regular | Orange | 100 | 5.000 |
| 304100 | 24G x 1" | 25 mm | Regular | Violet | 100 | 5.000 |
| 300700 | 23G x 1 1/4" | 30 mm | Thin | Blue | 100 | 5.000 |
| 300800 | 23G x 1" | 25 mm | Thin | Blue | 100 | 5.000 |
| 301000 | 22G x 1 1/2" | 40 mm | Thin | Black | 100 | 5.000 |
| 300900 | 22G x 1 1/4" | 30 mm | Thin | Black | 100 | 5.000 |
| 304727 | 22G x 1" | 25 mm | Thin | Black | 100 | 5.000 |
| 304432 | 21G x 1 1/2" | 40 mm | Thin | Green | 100 | 5.000 |
| 301156 | 21G x 1" | 25 mm | Thin | Green | 100 | 5.000 |
| 301300 | 20G x 1 1/2" | 40 mm | Thin | Yellow | 100 | 5.000 |
| 304827 | 20G x 1" | 25 mm | Thin | Yellow | 100 | 5.000 |
| 301500 | 19G x 1 1/2" | 40 mm | Thin | Ivory | 100 | 5.000 |
| 304622 | 18G x 1 1/2" | 40 mm | Thin | Pink | 100 | 5.000 |
| 301155 | 21G x 2" | 50 mm | Thin | Green | 100 | 4.000 |
| 300094 | 22G x 2" | 50mm | Regular | Black | 100 | 4.000 |
| 301900 | 18G x 2" | 50 mm | Regular | Pink | 100 | 4.000 |

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1.2 Certification

| BD REFERENCE | BD MANUFACTURER | ISO CERTIFICATION | CE MARKING | BD MANUFACTURING SITE |
|--|--|--|--|---|
| 300700, 300800, 300900, 301000, 301156, 301300, 301900, 304432, 304434, 304727, 304827 | Becton Dickinson S.A. - Crta. Mequinenza, s/n. 22520-Fraga (Huesca) Spain | AENOR - EN ISO 9001:2008 Certificate N° N° ER-0097/1994 AEMPS NB n°0318 EN ISO 13485:2013 Certificate N° 2015 05 0047EN | AEMPS N°0318 – Certificate n 95 06 0006 CP | Becton Dickinson S.A. - Crta. Mequinenza, s/n. 22520-Fraga (Huesca) Spain EN ISO 13485: 2013 Registration Number: 2015 05 0047 EN |
| 300094, 300300, 300600, 300635, 300637, 301500, 301700, 301900, 301750, 302200, 303800, 300400, 304100, 304300, 304622, 304000, 301155 | Becton Dickinson & Company Limited Donore Road Drogheda Co. Louth Ireland | NSAI – N° MD 19.1609 -EN ISO 13485:2012 | NSAI 050 – N° Q252.157 | Becton Dickinson S.A. - Crta. Mequinenza, s/n. 22520-Fraga (Huesca) Spain EN ISO 13485: 2013 Registration Number: 2015 05 0047 EN |



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1.3 Material

| COMPONENT | MATERIAL |
|---------------|--|
| Needle Hub | COLOR CODED POLYPROPYLENE |
| Needle Shield | POLYPROPYLENE |
| Bonding Agent | EPOXY |
| Needle | STAINLESS STEEL AISI 304 (Chromium 18-20%; Nickel 8-12%; Manganese 2%; Silicon 1%) |
| Lubricant | MEDICAL GRADE SILICONE OIL, <math><0.25 \text{ mg /cm}^2</math> |

1.4 Material of concern

Materials of concern are chemicals or substances that have been identified as having the potential to cause long term effects on humans or the environment.

| MATERIAL | COMMENT |
|-------------------------------------|--|
| Phthalates | No phthalates intentionally added. No DEHP, CAS number 117-81-1, EC number 204-211-0, intentionally added |
| Latex | The products do not contain natural latex. |
| Bisphenol A | Bisphenol A (CAS number 80-05-7, EC number 201-245-8) might be found in very low amount (at a concentration inferior to 5ppm) as a residue from the epoxy synthesis processing. Epoxy is used as needle bonding agent. |
| Substances of animal origin BSE/TSE | BD, Microlance devices utilize very small amounts of tallow or tallow derivatives (e.g. stearates in polymers). Per MEDDEV 2.4/1 Rev. 9 June 2010 and MEDDEV 2.11/1 Rev 2 January 2008, such substances are not considered as derivatives of animal tissues for the purpose of this rule which therefore does not apply. |
| Polyvinyl chloride (PVC) | The products do not contain polyvinyl chloride |

1.5 REACH information

BD maintains an active REACH compliance program and works closely with its supply base on an ongoing basis with a view to obtaining information on REACH Substances of Very High Concern (“SVHC”) through regular communication and exchange

1.6 Biocompatibility

BD Medical products comply with the requirements of the standard for biocompatibility of medical devices, ISO 10993-1 Biological Evaluation of Medical Devices - Part 1: Evaluation and Testing.

1.7 Sterilization

Ethylene Oxide Sterilization following *EN ISO 11135-1*. ETO residues are within applicable regulations.



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1.8 Shelf life

Shelf life 5 years. No special storage or transportation condition. Recommendations are to store in room temperature, in dry and warm place and not exposed to strong light.

1.9 Standards

| HARMONISED STANDARDS | |
|-----------------------------|--|
| EN 556-1:2002/ COR 2006 | Sterilisation of Medical Devices – requirements for medical devices to be labelled “sterile”. |
| EN 980: 2008 | Graphical Symbols for use in the labelling of medical devices. |
| BS EN 1041+A1: 2013 | Terminology, symbols and information provided with medical devices. Information supplied by the manufacturer with medical devices |
| EN 1707:1996 | Conical fittings with a 6 % (Luer) taper for syringes, needles and certain other medical equipment - Lock fittings |
| EN 20594-1:1993/AC:1996 | Conical fittings with a 6% (Luer) taper for syringes, needles and certain other medical equipment - Part 1: General requirements |
| EN ISO10993-1:2009/AC:2010 | Biological evaluation of medical devices - Part 1: Evaluation and testing |
| EN ISO 10993-3:2009 | Biological evaluation of medical devices - Part 3: Tests for genotoxicity, carcinogenicity and reproductive toxicity |
| EN ISO10993-4:2009 | Biological evaluation of medical devices - Part 4: Selection of tests for interactions with blood |
| EN ISO 10993-5:2009 | Biological evaluation of medical devices - Part 5: Tests for in vitro cytotoxicity |
| EN ISO 10993-7:2008/AC:2009 | Biological Evaluation of Medical Devices Part 7: Ethylene oxide sterilization residuals |
| EN ISO 10993-11:2009 | Biological evaluation of medical devices - Part 11: Tests for systemic toxicity |
| EN ISO10993-12:2012 | Biological evaluation of medical devices - Part 12: Sample preparation and reference materials |
| EN ISO 10993-13:2010 | Biological Evaluation of Medical Devices Part 13: Identification and quantification of degradation products from polymeric medical devices |
| EN ISO 10993-15:2009 | Biological evaluation of medical devices - Part 15: Identification and qualification of degradation products from metals and alloys |
| EN ISO 10993-17:2009 | Biological Evaluation of Medical Devices Part 17: Establishment of allowable limits for leachable substances |
| EN ISO 10993-18:2009 | Biological evaluation of medical devices - Part 18: Chemical characterisation of materials |
| EN ISO 11135-1:2007 | Sterilization of health care products - Ethylene oxide - Part 1: Requirements for development, validation and routine control of a sterilization process for medical devices |
| EN ISO 11138-2:2009 | Sterilization of health care products - Biological indicators - Part 2: Biological indicators for ethylene oxide sterilization processes |

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| | |
|---|--|
| EN ISO 11607-1:2009 | Packaging for terminally sterilized medical devices - Part 1: Requirements for materials, sterile barrier systems and packaging systems |
| EN ISO 11607-2:2006 | Packaging for terminally sterilized medical devices - Part 2: Validation requirements for forming, sealing and assembly processes |
| EN ISO 11737-1:2006/AC:2009 | Sterilization of medical devices - Microbiological methods - Part 1: Determination of a population of microorganisms on products |
| 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 ISO 13485:2012/AC:2012 and EN ISO 13485:2003 | Medical devices – Quality management Systems Requirements for Regulatory Purposes |
| EN ISO 14971:2012 | Medical Devices. Application of risk management to medical devices |
| EN 14155:2011 | Clinical investigation of medical devices for human subjects - Good clinical practice |
| NON HARMONISED STANDARD | |
| IS EN ISO 7884-1: 2016 | Sterile hypodermic needles For Single Use– Requirements and Test Methods |
| ISO 594-1:1993 | Conical fittings with a 6% (Luer) taper for syringes, needles and certain other medical equipment - Part 1: General requirements |
| ISO 594-2:1998 | Conical fittings with 6 % (Luer) taper for syringes, needles and certain other medical equipment -- Part 2: Lock fittings |
| ISO 9626: 1995 | Stainless steel needle tubing for the manufacture of medical devices |
| ISO 6009: 2016 | Hypodermic Needles for Single Use-Colour Coding for Identification |
| ISO 14644-1:1999 | Cleanrooms and associated controlled environments Part 1: Classification of air cleanliness |
| ISO 10993-2:2009 | Biological Evaluation of Medical Devices Part 2 |
| | |
| ISO10993-10:2009 | Biological evaluation of medical devices - Part 10: Tests for irritation and delayed-type hypersensitivity |
| ISO 15223-1:2012 | Graphical Symbols for use in the labelling of medical devices. |
| 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 |

1.10 Classification

Class IIa Medical Device under Rule 6, Annex IX of Medical Devices Directive 93/42/EEC as amended

1.11 GMDN code

GMDN code 59230

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1.12 Good Manufacturing practices

The entire manufacturing and testing processes are following the Good Manufacturing Practices as specified below

- Incoming raw materials are verified via material inspection and testing and our suppliers are approved via our vendor management system.
- In addition to the automatic on-line inspections, in-process inspections are performed in addition to final product testing to ensure compliance with approved specifications.
- The manufacturing and testing details of each batch of product are recorded on a batch record which is retained in accordance with our document control procedures
- BD operates a system of Internal and external audits to maintain compliance
- BD confirms that it will continue to adhere to relevant international standards in designing and manufacturing its products.
- BD reserves the right to use the internal change control procedure to change raw material suppliers and production process

1.13 Others

- (Material) Safety Data Sheets are not required for this product
- Certificate of Food Contact (*COMMISSION REGULATION (EU) No. 10/2011 of January 14th, 2011 concerning materials and plastic objects intended to get in touch with foodstuffs*) is not required as BD products are used for general purpose injection and aspiration of fluids from vials, ampoules and parts of the body below the surface of the skin.
- No separate Instruction for Use is available for these devices.

2. Packaging

2.1 Packaging material

LABELS: according to European Medical Device directive, multilingual

| | |
|---------------|-----------|
| Web packaging | POLYAMIDE |
| Box | PAPER |



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2.2 Example labeling

Example Unit pack label, SKU 300400, legal manufacturer Drogheda, from document DGW 924



Example Unit pack label, SKU 300800, legal manufacturer Fraga, from document DGW 920

