

TECHNICAL DATA SUMMARY (TDS)

MEDALLION SYRINGES (MSS)

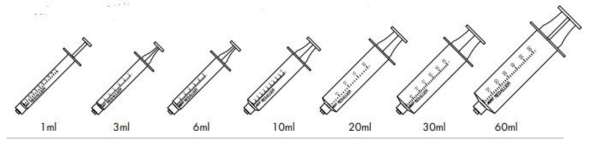
1. DEVICE DESCRIPTION

The Medallion® Series Syringes (MSS) are hypodermic 3-piece syringes constructed with clear barrels, color plungers, and a black piston seal. Available size options include 0.25, 1, 3, 6, 10, 20, 30, and 60 mL.

The syringe barrels are one-piece polycarbonate molded construction with finger grips in either a flat or "sword" style and connectors in a fixed male Luer, Toomey tip, or slip Luer style. Slip Luer styles are available with centric or eccentric nozzles. The syringe barrels are marked for accurate, clear and easy reading. The barrel may be customized with black or white printed text. The safety stop ring prevents inadvertent removal of the plunger from the syringe barrel. An attachable wing accessory is available for use with Merit's 1 mL and 3 mL Medallion® syringes. The wing accessory may be snapped onto the barrel of the syringe enabling a more secure grip when handled with wet, gloved hands. The Medallion® plungers are molded of Acrylonitrile-Butadiene-Styrene (ABS) and are fitted with a silicone rubber piston seal. Some configurations are also available with polycarbonate plungers.

The 60 mL Medallion® Syringe is constructed using the same materials and processes used in the manufacture of other Medallion® syringes except for the piston seal. The 60 mL Syringe utilizes an EPDM O-Ring seal instead of the silicone piston seal used in other syringes. The O-ring seal fits around the plunger head leaving plunger material in contact with fluid inside the syringe barrel.

The 0.25ml Medallion® Syringe is constructed with the same materials used in the manufacture of other Merit syringes except for the plunger, which is manufactured of glass-filled nylon. The 0.25mL Medallion Syringe is fitted with a silicone O-Ring piston seal. The O-ring seal fits around the plunger head leaving plunger material in contact with fluid inside the syringe barrel.



2. INTENDED USE / INDICATION FOR USE

The Merit Syringe is used to inject fluids into, or withdraw fluids from, the body.



3. Contraindications

There are no contraindications or warnings for this product.

4. INSTRUCTIONS FOR USE

The Instructions for Use and product labelling for the product have been formatted in multiple languages.

5. Classification:

a. European Community:

The classifications take into consideration of all rules based on the component with the highest classification. For devices claiming conformity to the Essential Requirements of the MDR, the following classification rules are used:

The product is not an active device, and none of the special rules apply.

Subjects	Applied Rules	Justification
Non-invasive devices	□ 1, ⊠ 2, □ 3, □ 4	See table below.
Invasive devices	□ 5, □ 6, □ 7, □ 8	Not an invasive device.
Active devices	□ 9, □ 10, □ 11, □ 12	Not an active device.
Special rules	□ 13, □ 14, □ 15, □ 16, □ 17, □ 18	None of the special rules apply.

Table 1: MDR Classification Rationale of Medallion

Component	Rationale	Rule	Class
Syringe b.	Non-invasive devices intended for channeling liquids for the purpose of eventual infusion, administration or introduction into the body	2	I – Sterile Measuring

b. United States Food and Drug Administration:

Class:

Product Code/CFR: FMF / 21 CFR 870.1650

510(k) Reference: K173601
Device Name: Piston Syringe

c. Canada:

Class: II (Rule 5)



License: 6072

d. GMDN and CND

GMDN Code: 47017 - General-purpose syringe, Single Use

CND Code: A0201 - Single Use Syringes

6. DEVICE SPECIFICATIONS

SPECIFICATION NAME	DESCRIPTION
Capacity	0.25 mL, 1 mL, 3 mL, 6 mL, 10 mL, 20 mL, 30 mL, 60 mL
Luer Connections	Fixed Male Luer & Male Slip Luer Per ISO 594-1/2 (or ISO 80369-7 testing currently underway)
Graduation Scale	Per ISO 7886-1

7. MATERIALS OF CONSTRUCTION

PRODUCT COMPONENT	MATERIAL
Barrel	Blue polycarbonate, clear polycarbonate
Plunger (1-60mL)	Acrylonitrile-Butadiene-Styrene (ABS) and Polycarbonate (PC)
Plunger (0.25mL)	Nylon
60 mL O-Ring	Ethylene Propylene Diene Methylene (EPDM) rubber
0.25 mL O-Ring	Silicone rubber
1-30 mL Tip	Silicone rubber
Lubricant	Medical grade silicone
Ink	Black or White U.V. Curable

PACKAGING COMPONENT	MATERIAL
Blister	Formed blister of 1073 Tyvek and EVA Trilayer film
Carton	Corrugate single wall

8. Packaging

Merit's syringes are packaged as sterile, single use devices. The barrier properties of the packaging assure that sterility is maintained. All units are packaged in a pre-formed blister tray and sealed with



medical grade Tyvek[™], or a Nylon pouch configuration. Medallion syringes are packaged 25 units per carton, 4 cartons per shipper.

Non-sterile syringe assemblies are packaged to facilitate incorporation into customized procedure packs which are provided to clinicians as sterile, single use devices. Product labeling of non-sterile devices clearly specifies the requirement for further processing (e.g. sterilization).

Certain bulk non-sterile syringe assemblies include labeling that specifies the validated sterilization parameters that repackers can use to leverage Merit's CE mark. These are triple poly-bagged and placed into cartons to protect the product and to reduce particulate contamination.





9. Labeling

Figure 2: Unit Label – Medallion

Medallion® Syringe





Figure 3: Carton Label - Medallion

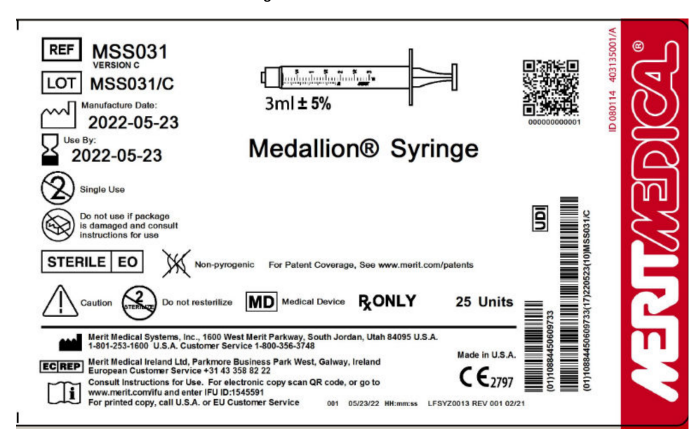
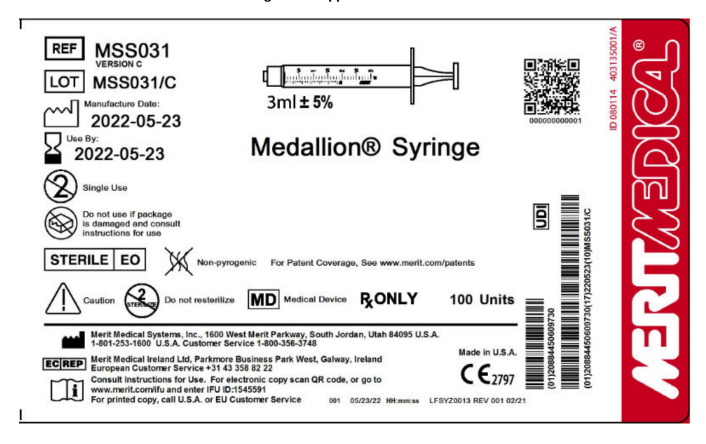




Figure 4: Shipper Label - Medallion



10. Compliance with Relevant Regulations and Technical Standards

EU:

Medical Device Regulation (MDR) of the European Union (Regulation (EU) 2017/745 of the European Parliament and of the Council of 5 April 2017 on medical devices). The applicable Certificate is MDR 716389.

US:

The product is legally marketed according to the requirements of the United States Food and Drug Administration in compliance with the premarket notification 510(k) process. The 510(k) clearance letter is K173601.

CAN:

Class: II (Rule 5) License: 6072



The qualification activities that were performed for the product were based on the below applicable standards:

DOCUMENT	TITLE	
Horizontal (General) Standards		
Council Directive 93/42/EEC	Medical Device Directive of the European Union	
Regulation (EU) 2017/745	Medical Device Regulation (MDR) of the European Union (Regulation (EU) 2017/745 of the European Parliament and of the Council of 5 April 2017 on medical devices)	
ISO 13485	Quality Systems – Medical Devices – Quality Management Systems. Requirements for Regulatory Purposes	
MEDDEV 2.1/5 (June 1998)	Medical Devices: Guidance Document: Medical Devices with a Measurement Function	
ISO 14971	Medical devices — Application of risk management to medical devices	
ISO 10993-1	Biological evaluation of medical devices Part 1: Evaluation and testing within a risk management process	
ISO 11607-1	Packaging for terminally sterilized medical devices – Part 1: Requirements for materials, sterile barrier systems, and packaging systems.	
ISO 11607-2	Packaging for terminally sterilized medical devices – Part 2: Validation requirements for forming, sealing and assembly processes	
ASTM D4169	Standard practice for performance testing of shipping containers and systems	
ASTM F1980	Standard guide for accelerated aging of sterile barrier systems for medical devices	
ASTM F2475	Standard Guide for Biocompatibility of Medical Device Packaging Materials	
ASTM F 2096	Standard Test Method for Detecting Gross Leaks in Medical Packaging by Internal Pressurization (Bubble Test)	
ASTM F 1929	Standard Test Method for Detecting Seal Leaks in porous Medical Packaging by Dye Penetration	
ASTM F88/F88M	Standard Test Method for Seal Strength of Flexible Barrier Materials.	
BS EN 980	Symbols for use in the labeling of medical devices	
EN 1041	Information supplied by the manufacturer of medical devices	
ISO 15223-1	Medical devices – Symbols to be used with medical device labels, labeling and information to be supplied – Part 1: General requirements	
ISO 11135-1	Sterilization of health care products Ethylene oxide Requirements for	



DOCUMENT	TITLE		
	development, validation and routine control of a sterilization process for medical devices		
ISO 11137-1	Sterilization of health care products Radiation Part 1: Requirements for development, validation and routine control of a sterilization process for medical devices		
EN 556-1	Sterilization of medical devices – Requirements for terminally-sterilized medical devices to be labeled "Sterile"		
USP	United States Pharmacopeias		
AAMI TIR 28	Product Adoption and process equivalency for ethylene oxide sterilization		
ANSI/AAMI ST72:2011	Bacterial Endotoxins – Test methods, routine monitoring, and alternatives to batch testing		
	Product Specific Standards		
ISO 7886-1	Sterile hypodermic syringes for single use – Part 1: Syringes for manual use		
ISO 594-1	Conical Fittings with 6% (Luer) Taper for Syringes, Needles and Certain Other Medical Equipment – Part 1: General Requirements		
ISO 594-2	Conical Fittings with 6% (Luer) Taper for Syringes, Needles and Certain Other Medical Equipment – Part 2: Lock Fittings		
ISO 80369-7	Small bore connectors for liquids and gases in healthcare applications – Part 7: Connectors for intravascular or hypodermic applications		

11. Storage and handling

Store under general warehouse conditions.

12. Biocompatibility

The Product meets ISO 10993-1 and the FDA guidance document: Use of International Standards ISO-10993-1, "Biological Evaluation of Medical Devices Part 1: Evaluation and Testing Within a Risk Management Process."

13. SHELF LIFE

The Products are labeled for a 5-year shelf life. Non-sterile units are labeled for a 5-year shelf life



14. STERILIZATION DETAILS

Merit Medical Systems, Inc. utilizes Ethylene Oxide (EO) sterilization for the products described in this Technical Data Summary. The sterilization cycle is validated according to European International Standard ISO 11135-1. These products meet ISO 10993-7 requirements for ethylene oxide sterilization residuals.

Merit also offers the option to Gamma sterilize some of the products described in this Technical Data Summary. The Gamma radiation process is validated using the Method VD_{max}^{25} – Substantiation of 25 kGy as the sterilization dose procedure from ISO 11137-2. Dose Audits are performed quarterly according to ISO 11137-2.

15. MATERIAL COMPLIANCE

EU REACH: To the best of our knowledge based on information currently available from our raw material suppliers as well as an assessment of the general material properties, the product does not contain substances listed on the ECHA Candidate List of Substances of Very High Concern (SVHC).

CMR and Endocrine-Disrupting Substances: An assessment of the raw materials used to manufacture the product in this summary with respect to carcinogenic, mutagenic or toxic to reproductive ('CMR') compounds, of category 1A or 1B, in accordance with Part 3 of Annex VI to Regulation (EC) No 1272 | 2008 of the European Parliament and of the Council, or substances having endocrine-disrupting properties was completed. No CMR compounds or substances having endocrine-disrupting properties are contained in the products that are the subject of this Technical Data Summary.

16. MANUFACTURING CONDITIONS / QUALITY CONTROL

Merit Medical Systems, Inc. manufactures the products in this summary in accordance with the United States Food and Drug Administration (FDA) Quality System Regulations, 21 CFR Part 820 *Quality Systems Regulations*.

Merit Medical Systems facilities used to manufacture the product in this summary are also certified to ISO 13485 Quality Management System.