DeltaVen® Closed System Catheter

Technical Summary Sheet

DEVICE CLASSIFICATION
Class IIb

DEVICE DESCRIPTION
Closed system safety IV catheter in polyurethane (PUR) with accessories.

GMDN
Code Number: 40601

CE MARK
CE 0123

PRODUCT DESCRIPTION
DeltaVen® is a radio-opaque polyurethane catheter equipped with a rubber septum inside which a stainless steel needle is inserted. The catheter is connected to a lateral extension equipped with a clamp (i) whose end section is connected to a fitting with a single or double Luer-lock complete with an air vent connection (f). The end connections can be equipped with a Luer-lock cap (e), with a needleless connector (d), with a three-way stopcock (g) which is also equipped with a needleless connector (h). The catheter is also equipped with a passive safety system to prevent needlestick injuries.

INDICATIONS
DeltaVen® is a catheter for peripheral venous and subcutaneous access that, in combination with other peripheral devices, allows the collection of blood samples and the administration of fluids. DeltaVen® is intended to be used for less than 30 days.

DeltaVen® is equipped with a passive system for the prevention of accidental needlestick injuries. During the initial insertion phase, the blood remains in the device facilitating the prevention of exposure to blood. The catheter can be used on any patient population while taking into account the vascular anatomy of the patient and of the adequacy of the procedure.

DeltaVen® catheters are suitable for use with pressure injectors (max. 330 psi) only if all accessories are removed. The recommended measurements for subcutaneous use are 26G, 24G and 22G with a 19 mm catheter length. DeltaVen® 26G: DO NOT USE FOR HIGH PRESSURE TREATMENTS.

PRECAUTIONS
Use of the device is for medical or paramedical competence.
- Read the instructions before use.
- Use protective gloves.
- Do not use if the individual packaging is damaged or open or if the product is past the “use by” date.
- Do not use if the device is incomplete or has been tampered with.
- The product must be used immediately after the packaging has been opened.
- Do not attempt to reinsert the entry needle in the rubber septum it was extracted from.
- Do not reinsert the needle while the catheter is located wholly or partly on site. It could break the catheter.
- Do not insert needles, sharp objects or faulty connectors in the needleless connector. It could cause leakages from the device. In case of an attempted insertion of a needle or a blunt cannula, replace the needleless connector.

Peripheral venous catheter with closed system (single and two-way) before removing the needle and the accessories.

Needle protection device after activation.
• Replace access without needle or needleless connector within a maximum time of 7 days or after exceeding 200 activations.

• If the device is equipped with a stopcock it is recommended that it be replaced after 72 hours or 24 hours if using concentrated solutions of glucose, lipids or blood products.

• If the catheter is in place and you want to replace the accessories, put the clamp in the closed position (OFF).

• Do not use needleless connector with an operating pressure exceeding 2 bar.

• Disposable device: do not sterilise and/or reuse in order to avoid compromising functionality or aid possible cross contamination with other patients.

• After use, dispose of as hospital waste.

• In case of incorrect transport and/or manipulation, the device or packaging could be subject to structural and/or functional damage.

• Do not use scissors at or near the insertion site.

• If pain, swelling around the catheter insertion area or the occurrence of edema/erythema or other local complications are experienced, replace the device and insert a new one in another site.

• Clean the device immediately after the administration of medicines or administration of small volumes of medicines.

• If pain, swelling around the catheter insertion area or the occurrence of edema/erythema or other local complications are experienced, replace the device and insert a new one in another site.

• Immediately remove any needle that has no coating, always keeping the tip away from your body and fingers.

• Do not expose to heat or direct sunlight.

• If the catheter is incorrectly inserted under the skin, remove it and never try to re-insert the needle into the catheter when it was extracted partially or totally.

• If during insertion under the skin blood appears in the needle mounting unit or in the catheter after withdrawing the needle, discard the device and insert in another place.

• If you want to administer a drug bolus different from what is being administered subcutaneously in a continuous mode, you must insert a new catheter to avoid incompatibilities between drugs.

If the device is used at a high pressure or with injectors:

• Connect directly to the pressure infusion system with the end Luer-lock connector of the device.

• Remove all the accessories connected to the device and replace them with a Luer-lock cap where necessary.

• Always check the patency of the device before use.

• Never exceed the maximum pressure of 23 bar (330 psi).

• Consider the internal residual volume of the device in the case of administration of small volumes of medicines.

• If pain, swelling around the catheter insertion area or the occurrence of edema/erythema or other local complications are experienced, replace the device and insert a new one in another site.

• Clean the device immediately after the administration of medicines or biological fluids.

• Immediately remove any needle that has no coating, always keeping the tip away from your body and fingers.

• Do not expose to heat or direct sunlight.

• If the catheter is incorrectly inserted under the skin, remove it and never try to re-insert the needle into the catheter when it was extracted partially or totally.

• If during insertion under the skin blood appears in the needle mounting unit or in the catheter after withdrawing the needle, discard the device and insert in another place.

• If you want to administer a drug bolus different from what is being administered subcutaneously in a continuous mode, you must insert a new catheter to avoid incompatibilities between drugs.

PRODUCT COMPONENTS
Non-pyrogenic, sterile, single-use device. The device is not made from natural latex and phthalates.

DeltaVen® is a radio-opaque polyurethane catheter equipped with a rubber septum inside which a stainless steel needle is inserted. The catheter is connected to a lateral extension equipped with a clamp (l) whose end section is connected to a fitting with a single or double Luer-lock complete with an air vent connection (f). The end connections can be equipped with a Luer-lock cap (e), with a needleless connector (d), with a three-way stopcock (g) which is also equipped with a needleless connector (h). The catheter is also equipped with a passive safety system to prevent needlestick injuries.

COMPONENT COMPOSITION
Polyurethane catheter with three radio-opaque strips enclosed in the wall containing BaSO4.

• AISI304 stainless steel needle.

• Polycarbonate catheter hub.

• Silicone rubber septum needle seal.

• Butterfly with polypropylene body and thermoplastic synthetic rubber wings.

• Polyurethane extension line DEHP free.

• ABS/Polypropylene colored Luer-lock connection.

• Styrene butadiene copolymer air vent with high density polyethylene hydrophobic filter.

• Polypropylene Luer-lock cap.

• Polycarbonate stopcock with silicone rubber needleless connector. Necessary connector with polyester/polycarbonate hub and silicone rubber core.

MANUFACTURING SITE NAME AND ADDRESS
Delta Med SpA
Via G. Rossa, 20-44019
Viadana (Mn) - Italy
0375 785915

COUNTRY OF ORIGIN
Italy

ANIMAL ORIGIN
None of the devices listed on this product family incorporate any substances that are derived from animal tissues as per Commission Regulation (EU) 722/2012.

STERILIZATION METHOD
Ethylene Oxide

SHELF-LIFE
59 months

LABELLING AND PACKAGING

For more information visit our website at www.smiths-medical.com

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