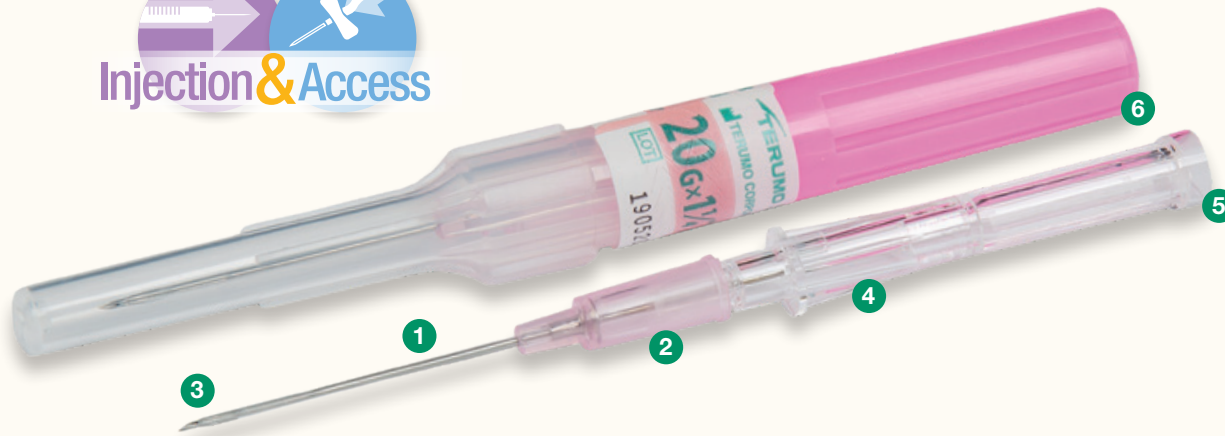


HOSPITAL CARE

Surflash™

I.V. Catheter
Product Information



When using the Surflash™ catheter, the health care provider has a visual confirmation of correct insertion of the catheter in the vein through the Surflash™ sytem.

Description	Raw Materials
1 Catheter	Polyurethane + BaSO ₄
2 Catheter hub Caulking pin	Polypropylene + master batch Stainless steel
3 Cannula (internal needle)	Stainless steel
4 Needle hub	Polycarbonate
5 Air filter	Hydrophobic filtering membrane/Polyester chlorinated PVC
6 Case with coloured cap	Polypropylene

Does not contain components made of natural rubber latex.

Sterilization

With ethylene oxide to a Sterility Assurance Level of at least 10⁻⁶

Storage

- Shelf life of 5 years
- The batch number and expiry date are printed on the label of the unit pack
- Do not store at extreme temperature and humidity

Equipped with a placement confirmation

Your catheters of choice for a wide range of I.V. applications

Product Range

Product Code	Colour	External Catheter Ø		Internal Catheter Ø	Catheter Length	Flow Rate ml/min.
SR*FF2419	Yellow	24G	0.67 mm	0.47 mm	19 mm	15
SR*FF2225	Blue	22G	0.85 mm	0.60 mm	25 mm	38
SR*FF2051	Pink	20G	1.10 mm	0.80 mm	51 mm	59
SR*FF2032	Pink	20G	1.10 mm	0.80 mm	32 mm	67
SR*FF1851	Green	18G	1.30 mm	0.95 mm	51 mm	88
SR*FF1832	Green	18G	1.30 mm	0.95 mm	32 mm	99
SR*FF1651	Grey	16G	1.70 mm	1.30 mm	51 mm	184
SR*FF1464	Orange	14G	2.17 mm	1.73 mm	64 mm	282



Precautions

- Do not use if unit package is damaged
- Do not re-use
- Do not attempt to re-insert a partially or completely withdrawn needle
- Safely dispose of all contaminated materials according to facility protocol observing appropriate local guidelines for biohazard disposal.
- Catheter should be changed according to CDC guidelines or facility policy
- Due to the increased tendency for pediatric patients (in comparison to adult patients) to have a stronger reaction to physiologic stress that may be associated with certain procedures, appropriate precautionary measures should be taken when this device is used on a child.

Always consult the product label and instructions for use (IFU) for a complete overview of warnings, cautions and/or precautions prior to actual use.

Packaging

Unit Pack Hard Case

Unit Box 50 pieces

Shipping Carton

200 pieces

Gross Weight 1.08 kg

Dimensions

266 x 690 x 236 mm

Regulatory Status

CE Certification – Class IIa medical device

Notified body

TÜV Rheinland 0197

Manufacturer

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EC Representative

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This range complements the wider Terumo product offering including a comprehensive portfolio of both safety and conventional I.V. products which can be found at our web page on www.terumo-europe.com

Your Terumo partner



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