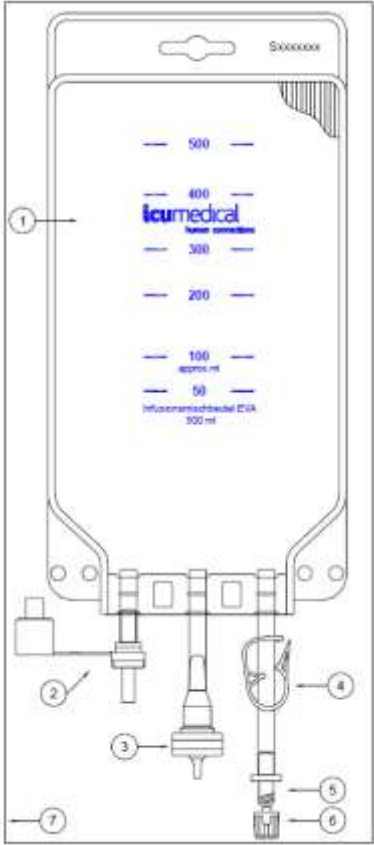


PRODUCT TECHNICAL DATA SHEET	
ITEM REFERENCE	SN2050
LEGAL MANUFACTURER	ICU Medical, Inc. 951 Calle Amanecer, San Clemente, CA 92673, USA
ASSEMBLY SITE	<ul style="list-style-type: none"> <li>ICU Medical, Inc., 4455 Atherton Drive Salt Lake City, UT 84123</li> <li>ICU Medical S. de R.L. de C.V., Avenida Cuarza No. 250 Colonia Rancho, Santa Clara, Manáedero Ensenada, Baja California, Mexico 22790</li> </ul>
ITEM DESCRIPTION	INFUSION BAG EVA 500ml CLAMP ON/OFF
INTENDED USE	Parenteral / Enteral Solution Bag
	
DEVICE GENERIC INFORMATION	Infusion bag EVA 500 ml with clamp.



**ICU MEDICAL EUROPE srl**  
A socio unico

**Società soggetta ad attività di direzione e coordinamento di ICU Medical Inc. USA**

Sede legale e operativa: Roncanova di Gazzo Veronese, Via Martiri della Libertà, 1/A, 37060 Roncanova (VR)

Registro Imprese di Verona: 03237150234 - REA di Verona: 320421 - CF e P.IVA: IT 03237150234

Tel. +39 0442 570106 - Fax +39 0442 570109

<b>LIST OF COMPONENTS</b>	<p>Infusion bag 500ml (EVA)</p> <p>Infusion point Ø 4.8mm with cap (ABS, Polypropylene)</p> <p>Infusion Clousure break open</p> <p>Pinch clamp medium, Roberts Model, white color</p> <p>Female Luer Lock (ABS) with vented cap (LDPE)</p>
<b>PACKING AND PACKAGING</b>	<p>Individual packaged product using pouch/blister in medical paper and film.</p> <p>Cardboard boxes.</p> <p><b>Indivisible box composed by: 40 pcs.</b></p> <p>Packing and packaging according to UNI EN ISO 11607/1-2.</p>
<b>STERILIZATION</b>	<b>ETO:</b> v alidated method according to UNI EN ISO 11135
<b>EXPIRY – SHIELD LIFE</b>	<p>5 Year (f rom sterilization date) if undamaged package and properly stored</p> <p>Do not re-use  , Reuse negatively impacts performance / sterility potentially resulting in product failure / contamination.</p>
<b>BIOCOMPATIBILITY</b>	All the material are biocompatible according to ISO 10993
<b>LABELS</b>	Labels are applied on the individual bags and on the outside of the cardboard box. Product identification data according to Directive 93/42/EEC point 13.3 and the specific technical standards
<b>INSTRUCTION FOR USE</b>	Contained in each sales package (according to EN 1041 and further modifies and integrations)
<b>PRODUCTION ENVIROMENT</b>	Controlled production environments. Microbiological and particle controls are performed on environmental air and surfaces (UNI EN ISO 14644/1-2).
<b>TRACEABILITY</b>	ICU Medical guarantees full traceability of all the components used in the production of its devices
<b>DISPOSAL</b>	The user must dispose the device according to hospital disposal policy.
<b>STORAGE</b>	Store in a dry and clean place.
<b>WARNINGS</b>	Use aseptic techniques. Single-use only – Do not re-sterilize.
<b>PRODUCTION CONTROLS</b>	<ul style="list-style-type: none"> <li>Each component is inspected during acceptance (visual inspections, dimensional checks and functional tests) to verify conformity with the requirements according to ICU Medical internal quality procedures</li> <li>During product production and release specific tests are performed according to ICU Medical internal quality procedures.</li> </ul>
<b>QUALITY SYSTEM AND PRODUCT CERTIFICATION</b>	<p><b>Quality System in compliance with I.S.EN ISO 13485:2012</b></p> <p>➤ <b>Certificate</b> Number MD19.4496</p> <p><b>Notified body:</b> NSAI (National Standards Authority of Ireland)</p>
	<p><b>Product certification:</b> in compliance with Directive 93/42/EEC (Annex II) and further modifies and integrations.</p> <p><b>CE Certificate:</b> Number 252.884 </p> <p><b>Notified Body:</b> NSAI (National Standards Authority of Ireland)</p> <p><b>Classification:</b> Class IIa (Annex IX Directive 93/42/EEC).</p>

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