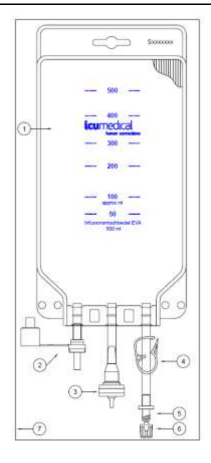


PRODUCT TECHNICAL DATA SHEET		
ITEM REFERENCE	SN2050	
LEGAL MANUFACTURER	ICU Medical, Inc. 951 Calle Amanecer, San Clemente, CA 92673, USA	
ASSEMBLY SITE	 ICU Medical, Inc., 4455 Atherton Drive Salt Lake City, UT 84123 ICU Medical S. de R.L. de C.V., Avenida Cuarza No. 250 Colonia Rancho, Santa Clara, Manaedero Ensenada, Baja Califomia, Mexico 22790 	
ITEM DESCRIPTION	INFUSION BAG EVA 500ml CLAMP ON/OFF	
INTENDED USE	Parenteral / Enteral Solution Bag	



DEVICE GENERIC Inf usion bag EVA 500 ml with clamp.

INFORMATION

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ICU MEDICAL EUROPE srl A socio unico

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



	Influeing hop FOOm (FVA)
LIST OF COMPONENTS	Infusion bag 500ml (EVA) Infusion point Ø 4.8mm with cap (ABS, Polypropylene) Infusion Clousure break open Pinch clamp medium, Roberts Model, white color Female Luer Lock (ABS) with vented cap (LDPE)
PACKING AND PACKAGIN	Individual packaged product using pouch/blister in medical paper and film. Cardboard boxes. Indivisible box composed by: 40 pcs. Packing and packaging according to UNI EN ISO 11607/1-2.
STERILIZATION	ETO: v alidated method according to UNI EN ISO 11135
EXPIRY – SHIELD LIFE	5 Year (from sterilization date) if undamaged package and properly stored Do not re-use , Reuse negatively impacts performance / sterility potentially resulting in product failure / contamination.
BIOCOMPATIBILITY	All the material are biocompatible according to ISO 10993
LABELS	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
INSTRUCTION FOR USE	Contained in each sales package (according to EN 1041 and further modifies and integrations)
PRODUCTION ENVIROMENT	Controlled production environments. Microbiological and particle controls are performed on environmental air and surfaces (UNI EN ISO 14644/1-2).
TRACEABILITY	ICU Medical guarantees full traceability of all the components used in the production of its devices
DISPOSAL	The user must dispose the device according to hospital disposal policy.
STORAGE	Store in a dry and clean place.
WARNINGS	Use aseptic techniques. Single-use only – Do not resterilize.
PRODUCTION CONTROLS	 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 During product production and release specific tests are performed according to ICU Medical internal quality procedures.
QUALITY SYSTEM AND PRODUCT CERTIFICATION	Quality System in compliance with I.S.EN ISO 13485:2012 > Certificate Number MD19.4496 Notified body: NSAI (National Standards Authority of Ireland) Product certification: in compliance with Directive 93/42/EEC (Annex II) and further modifies and integrations. CE Certificate: Number 252.884 Notified Body: NSAI (National Standards Authority of Ireland) Classification: Class IIa (Annex IX Directive 93/42/EEC).

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