



Product PN GSS001A02S GSS001B02S GSS002A02S GSS002B02S

Mod. 984e-ext

Description HI-FLO Extension Set Adult 0.2/1.2 µm Y-Infusion connector

Rev. 06

HI-FLO Extension Set Adult with 0.2/1.2 µm Filter and Y-injection



PRODUCT
DESCRIPTION

Infusion set for infusion for intravascular applications.

The device is sterile and can be used in combination with an infusion set or an infusion device (pump or syringe).

The connections are a female luer lock with vented cap inlet and a male luer lock with cap ('A' model) or male luer lock with ring and protective cap ('B' model), compliant with ISO 80369-7.

The set is provided with a vented Speedflow® Adult filter, with an hydrophilic PES membrane with 0.2/1.2 μm pore size for particles retention and a hydrophobic PTFE membrane with 0,03 μm pore size.

The set is provided with a y-injection site, for administration of additional drugs by means of a syringe using the same infusion line.

The approximate total length of the set is 630 mm.

The set includes the following components:

- Female luer lock with vented cap inlet;
- Y-Injection point;
- Speedflow Adult 0.2/1.2 µm IV Filter;
- Male luer lock with cap/with ring and protective cap outlet.

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GSS002A02S
GSS001B02S
0.2 μm with male luer lock with cap outlet
1.2 μm with male luer lock with cap outlet
0.2 μm with male luer lock with ring and p

1.2 µm with male luer lock with cap outlet0.2 µm with male luer lock with ring and protective cap outlet1.2 µm with male luer lock with ring and protective cap outlet

MANUFACTURER NAME

GVS SpA Via Roma, 50

GSS002B02S

40069 Zola Predosa (BO) - ITALY

Phone: +39.051.6176311 – Fax: +39.051.6176200 e-mail: <u>gvs@gvs.it</u> – website: <u>www.gvs.com</u>

INTENDED USE / APPLICATION

Infusion of Drugs (including chemotherapy), Anesthetics, Nutritional Solutions and Saline Solutions for intravascular applications.

The set can be connected via Luer Locks to all commercially available infusion administration sets compliant with ISO 80369-7.

The set can be used either for gravity or in combination with an infusion set or an infusion device (pump or syringe). The set with 0.2 µm filter is intended specifically to eliminate air bubbles and to retain particles and bacteria, while the set with 1.2 µm filter is intended specifically to eliminate air bubbles and to retain fungi and microorganisms.

The set is a single-use device that can be used applications that last:

- Devices with 0.2 µm filter: up to 96 hours;

The set has to be disposed after each therapy.

Sets with 0.2 µm filter cannot be used for infusion of hyper alimentation liquids, lipids, blood and/or blood derivatives. Sets with 1.2 µm filter cannot be used for infusion of blood and/or blood derivatives.

The device should only be supervised and used by qualified healthcare personnel.

CLASS OF THE PRODUCT

Class IIa – sterile – single use

Rule 2 and 3 Annex VIII 2017/472/UE

EMDN A03020101

LOW PRESSURE EXTENSION LINES

MATERIALS

Filter media: PES Hydrophilic membrane / PTFE Hydrophobic membrane

Filter housing: MBS

Tubes: PVC (DEHP Free)





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Y-injection: SEBS / MABS Luer Lock: PVC / MABS Caps: HDPE

Regulatory Compliance:

- Biocompatibility according to ISO 10993-1
- Rohs directive 2011/65/EU
- DEHP plasticizer Free
- Latex free
- Reach 1907/2006/CE (hazardous substances regulation)
- Reg. 1272/2008/CE (classification, labelling and packaging of substances and mixtures)

PRODUCT CHARACTERISTICS

Physical/Mechanical:

Approximate total length: 630 mm
Total internal volume of the set: 11 ml
Weight (approx.): 14.4 g

Input/output connectors: Female luer lock with vented cap inlet compliant with ISO 80369-7

'A' model: Male luer lock with cap outlet compliant with ISO 80369-7 'B' model: Male luer lock with ring and protective cap outlet compliant

with ISO 80369-7 From 5 °C to 40 °C

Operating temperature Range: From 5 °C to 40 °C Storage temperature Range: From 0 °C to 40 °C Maximum applicable pressure: 3.2 bar

Biological:

Biocompatibility: Compliant with ISO 10993-1

Features:

Type of administration: gravity / pressure

Duration of the application: 0.2 μm up to 96 hours
1.2 μm up to 24 hours

Filter: Speedflow Adult 0.2/1.2 μm vented

Filter pore size: 0.2 µm
Filter internal volume: < 2.4 ml

Filter media: hydrophilic PES membrane and hydrophobic PTFE membrane

Flow Regulator: No
Drip Chamber: No
Roller: No
Y-injection site: Yes
Clamp: No

Tubing: PVC tube – inner Ø 3.0 mm / outer Ø 4.1 mm

PRODUCT SHELF LIFE 5 years

STERILIZATION Sterile: Yes – EtO

Suitable for Sterilization/Re-sterilization: No

APPLICABLE STANDARDS AND REGULATIONS **Product Certification:**

CE mark

Applicable Standards and Technical Regulations:

EN 556-1 Sterilization of medical devices — Requirements for medical devices to be designated "STERILE"

Part 1: Requirements for terminally sterilized medical devices

EN ISO 8536-4 Infusion equipment for medical use — Part 4: Infusion sets for single use, gravity feed

EN ISO 8536-9 Infusion equipment for medical use — Part 9: Fluid lines for single use with pressure infusion

equipment

EN ISO 8536-11 Infusion equipment for medical use — Part 11: Infusion filters for single use with pressure infusion

equipment

EN ISO 10993-1 Biological evaluation of medical devices — Part 1: Evaluation and testing within a risk

management process





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EN ISO 10993-4	Biological evaluation of medical devices — Part 4: Selection of tests for interactions with blood				
EN ISO 10993-5	Biological evaluation of medical devices — Part 5: Tests for in vitro cytotoxicity				
EN ISO 10993-7	Biological evaluation of medical devices — Part 7: Ethylene oxide sterilization residuals				
EN ISO 10993-10	Biological evaluation of medical devices — Part 10: Tests for irritation and skin sensitization				
EN ISO 10993-11	Biological evaluation of medical devices — Part 11: Tests for systemic toxicity				
EN ISO 10993-18	Biological evaluation of medical devices — Part 18: Chemical characterization of materials				
EN ISO 10993-23	Biological evaluation of medical devices — Part 23: Tests for irritation				
EN ISO 11135	Sterilization of health-care products — Ethylene oxide — Requirements for the development, validation and routine control of a sterilization process for medical devices				
EN ISO 11607-1	Packaging for terminally sterilized medical devices — Part 1: Requirements for materials, sterile barrier systems and packaging systems				
EN ISO 11607-2	Packaging for terminally sterilized medical devices — Part 2: Validation requirements for forming, sealing and assembly processes				
EN ISO 13485	Medical devices — Quality management systems — Requirements for regulatory purposes				
EN ISO 14971	Medical devices — Application of risk management to medical devices				
EN ISO 15223-1 Medical devices — Symbols to be used with medical device labels, labelling and inform supplied - Part 1: General requirements EN ISO 20417 Medical devices — Information to be supplied by the manufacturer					
					IEC 62366-1 Medical devices — Part 1: Application of usability engineering to m
EN ISO 80369-1	-1 Small-bore connectors for liquids and gases in healthcare applications — Part 1: General requirements				
EN ISO 80369-7	mall-bore connectors for liquids and gases in healthcare applications — Part 7: Connectors for travascular or hypodermic applications				
EN ISO 80369-20	Small-bore connectors for liquids and gases in healthcare applications — Part 20: Common test methods				
Available languages: English / Italian / German / French / Spanish					
Primary Packaging	l:				
,	Devices are individually packed and label in medical paper bags. Bag Size: 130 X 200 mm				
Secondary Packaging:					
Coomany rankag	Bags are placed inside a microperforated sack.				
Tertiary Packaging: Carton Box.					
	Box Size: 60 x 40 x 20 cm				
	Box Weight: 2.1 Kg Devices per box: 100				
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ISO 9001:2015 ISO 13485:2016 CE Certificate (201	7/745/UE)				
	EN ISO 10993-5 EN ISO 10993-7 EN ISO 10993-10 EN ISO 10993-11 EN ISO 10993-11 EN ISO 10993-23 EN ISO 10993-23 EN ISO 11135 EN ISO 11607-1 EN ISO 11607-2 EN ISO 14971 EN ISO 14971 EN ISO 15223-1 EN ISO 20417 IEC 62366-1 EN ISO 80369-7 EN ISO 80369-10 ISO 80369-10 ISO 9001:2015 ISO 9001:2015 ISO 9001:2015 ISO 9001:2015				



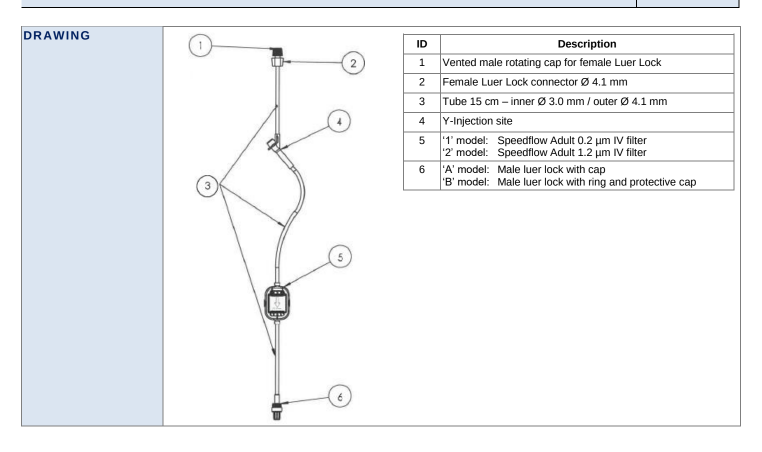
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REVISIONS AND APPROVALS:

DATE	REV.	REASON FOR CHANGE		VERIFIED BY: (name/function/signature)	APPROVED BY: (name/function/signature)
26/02/2023	00	First issue	Barbara Palmieri RA Specialist Bouloulalis	Barbara Finessi QA Manage Follow fun	Luca Zanini VP Healthcare and Lifesience