

# InVision-Plus CS® Chlorhexidine + Silver Engineering with Neutral Advantage Technology

Caution: Federal (USA) law restricts this device to sale by or on the order of a physician

## Indications for Use

The RyMed Technologies InVision-Plus CS® with Neutral Advantage Technology is a needleless IV connector system and is intended for single patient use in intravenous and blood administration sets without need for needles, thus eliminating the potential for needle-stick injuries during use.

The InVision-Plus CS® is antibacterial, containing silver and chlorhexidine, which are intended to inhibit the growth of microorganisms on the treated surfaces of the device, which include the septum and the fluid path.

*In vitro* testing of the InVision-Plus CS® has been shown to be effective for up to 7 days against the following microorganisms: Acinetobacter baumannii, Methicillin-resistant Staphylococcus aureus (MRSA), Staphylococcus aureus, Escherichia coli, Candida albicans, Klebsiella pneumoniae, Pseudomonas aeruginosa, and Staphylococcus epidermidis.

Organism	Log Reduction			
	Day 1	Day 3	Day 4	Day 7
MRSA (clinical isolate)	3.2	5.8	5.5	7.0
E. coli (ATCC #8739)	4.1	4.1	5.2	4.9
S. aureus (ACCT #6538)	3.2	5.2	4.2	4.2
P. aeruginosa (ACCT #9027)	4.3	4.9	4.3	4.5
A. baumannii (ATCC #19606)	3.8	3.8	4.5	5.4
S. epidermidis (clinical isolate)	4.0	5.4	4.0	6.0
K. pneumoniae (ATCC #4352)	5.0	4.2	5.5	5.1
C. albicans (ATCC #38245)	2.6	3.5	3.3	2.9

The subject device is not intended to treat existing infections. The device is not intended to have any effect on contaminated infusion solutions. Correlation between *in vitro* antibacterial activity and any clinical effectiveness has not been tested. The InVision-Plus CS® may be used with low pressure power injectors having a maximum pressure of 325 psi and a maximum flow rate of 10 mL/sec.

**Device Description:** The InVision-Plus CS® is accessed by a standard male-luer syringe or I.V. set connector. A male luer-lock connector should be used with the InVision-Plus CS® for a secure connection. The InVision-Plus CS® is bi-directional and luer-locking. The InVision-Plus CS® will automatically close when the syringe or male-luer I.V. set connector is removed. The InVision-Plus CS® may be used with low pressure power injectors having a maximum pressure of 325 psi and a maximum flow rate of 10mL/sec. Connect only with other devices rated for high pressure with a luer lock connection when performing power injections. The subject device should be changed after each use with a low pressure power injector.

The Neutral Advantage Technology features include a smooth swabbable septum surface, septum seal integrity with no gaps or openings, patented double microbial barrier, straight-through fluid pathway, zero dead space, zero fluid displacement, 0.027mL priming volume, 100% effective blood clearing, saline-only flush option and no clamping sequence or positive pressure syringe technique required.

**Contraindications:** The InVision-Plus CS® is contraindicated for patients with hypersensitivity to chlorhexidine or silver.

**How Supplied:** The InVision-Plus CS® is supplied as a standalone device or as part of extension and IV administration sets.

**Precautions:** Always use aseptic technique. The InVision-Plus CS® may not be compatible with every male-luer connector. Some I.V. sets/syringes utilize a male-luer connector with a lumen opening diameter of 0.063" or less which may cause damage to the InVision-Plus CS® resulting in septum damage, reduction or loss of fluid flow. DO NOT use any male-luer device unless certain that the male-luer connector opening is 0.064" or larger. Do not use needles with connector. Do not use luer-lock end caps on connector. Do not swab with Chlorhexidine Gluconate (CHG), use 70% Isopropyl Alcohol only. Flush InVision-Plus CS® after each use per institutional protocol using normal saline, or normal saline and heparin. Change per CDC guidelines or per validated facility protocol. In accordance with INS Standards, when using a Huber needle with a Y-Site or bifurcated tubing, use the access site closest to the patient as primary access. Clamp the alternate extension tubing when not in use. The CDC recommends that tubing used to administer lipid emulsions and TNA solutions should be changed every 24 hours. The InVision-Plus CS® and extension set should also be changed at this time.

## Instructions for Use - Needleless IV connectors


1. Wash hands thoroughly with bactericidal soap before each procedure. Use gloves, if required by your institutional protocol.
2. Inspect InVision-Plus CS®. Discard the set if end caps are missing or loose in package.
3. RYM-7000: Peel back lid on the "Touch-Free" Package Container. Do not touch the tip of the InVision-Plus CS®. Attach InVision-Plus CS® to the desired vascular access device by rotating the "Touch-Free" Package Container in a clockwise direction until secure. Pull back the Package Container until clear of the port and dispose.  
RYM-7001: Peel back lid on the Blister Package. Remove the male-luer cover. Do not touch the tip of the InVision-Plus CS®. Attach the InVision-Plus CS® to the desired vascular access device by rotating the InVision-Plus CS® in a clockwise direction until secure.
4. Remove protective coverings as assembly progresses.
5. To access the InVision-Plus CS® swab the septum and thread area with 70% Isopropyl Alcohol.
6. Position the syringe or I.V. set male-luer connector tip on the InVision-Plus CS® septum.
7. Push forward, rotate syringe or I.V. set male luer-lock connector clockwise until secure.
8. To disconnect the syringe or I.V. set male-luer connector, rotate counterclockwise.

## Instructions for Use - Specific to InVision-Plus CS® Extension Sets and Administration Sets

1. Open clamp, then prime to expel air from set.
2. Attach set to vascular access device.
3. Ensure that all connections are tightened and secure before use.

InVision-Plus CS® is compatible with magnetic resonance imaging (MRI) procedures

 Radiation Sterilized

 Do Not Reuse

Do Not Re-Sterilize

Rx Only

 Expiration Date

 Consult Instructions for Use

Not made with DEHP or natural rubber latex

Non-Pyrogenic Fluid Pathway

Manufactured for RyMed Technologies, LLC 6000 W. William Cannon Dr. B300 Austin, TX 78749 USA 1-512-301-7334

InVision-Plus CS® is a registered trademark of RyMed Technologies, LLC  
Patents: #6,113,068; #6,299,131; #6,994,315; #7,530,546B2; #8,096,525B2 and other patents pending  
RyMed Technologies, LLC 2016

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