

ProSeal™ Priming Cap

REF 423150

Priming Volume 0.17 ml

Packaging

100 pcs / Inner Box

400 pcs / Outer Carton

NOT MADE with NATURAL RUBBER LATEX or DEHP



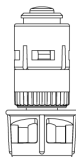
EC REP

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Indications for Use:

The ProSeal Closed System drug Transfer Device (CSTD) mechanically prohibits environmental contaminants from entering the system and the escape of drug or vapor concentrations from the system, thereby minimizing individual and environmental exposure to drug vapor, aerosols, and spills. The ProSeal system also prevents the introduction of microbial contaminants into the drug or fluid path for up to 168 hours or 7 days.

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STERILE LEO



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ProSeal Priming Cap enables the priming of closed system infusion lines. When connected to a ProSeal Injector (Syringe Adaptor) Ref 421010 at the distal end of infusion line, fluid can be primed in a closed system. The hydrophobic filter on the priming cap could stop the priming fluid flow once infusion line is fully primed.

The ProSeal Priming Cap and all its corresponding interface membranes exhibit a dry connection with the communicating surfaces in a fluid transfer. The use of this component and its appropriate ProSeal CSTD connecting component reduces the risk of microbial ingress for up to 168 hours or 7 days.

DIRECTIONS - Use Aseptic Technique

1. Prior to priming, swab the top of ProSeal Priming Cap with 70% isopropyl alcohol (15 seconds) and allow to dry (approximately 30 secs). Drying time is dependent on temperature, humidity, and ventilation area.
2. Attach ProSeal Priming Cap to the ProSeal Injector (Syringe Adaptor) Ref 421010 on the infusion line or syringe to be primed.
3. Start the priming. The priming fluid flow will be stopped automatically by the hydrophobic filter on priming cap once the infusion line is fully primed.
5. Remove used ProSeal Priming Cap and discard in accordance to disposal procedures for biohazardous materials of your facility.

Contraindications

The device is contraindicated whenever:

- The drug to be prepared is contraindicated to polycarbonate, ABS, stainless steel, polypropylene, silicone and polyisoprene.

Warnings

- Use accepted IV and pharmacy practice.
- The performance of the self-sealing membrane of the device is reduced after multiple perforations.
- Do not re-use to avoid contamination.

Precautions

- Do not use when caps and/or components are loose.
- The device is sterile unless packaging is damaged. Do not use when packaging is damaged.