Universal Funnel Adapter - Luer Lock

Medical device

Do not re-use

Non-pyrogenic
Non made with natural rubber latex

Do not resterilize

Consult instructions for use/electronic instructions for use

Sanitized using ethylene oxide
Single sterile barrier system

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UreSil is a registered trademark of UreSil, LLC.
Instructions for use include English.

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**UreSil ® Universal Funnel Adapter - Luer Lock**

**Medical device**

**INTENDED USE:** For connecting a percutaneous drainage catheter to a drainage bag or system. This product is intended for short-term use (less than 30 days).

**INTENDED PATIENT POPULATION:** This product is intended to be used with patients with placed drainage catheters.

**INTENDED USERS:** This product is intended only for use by trained clinicians.

**CLINICAL BENEFITS:** Work as an accessory in conjunction with a percutaneous drainage catheter and a Gravity Drainage bag or system to remove fluids using gravity or in conjunction with a percutaneous drainage catheter and a Suction Drainage bag or system to remove fluids using active suction or gravity.

**PERFORMANCE CHARACTERISTIC:** UFA-LL is comprised of a tapered funnel and a female luer connector and is used with funneled catheters, for example, UreSil Ulti-flo Large Bore non-locking catheters (REF LBNL-series), to connect to a Gravity Drainage or Suction Drainage bag or system with standard luer.
INTENDED FOR USE WITH: This accessory for a medical device is intended for use with funneled catheters, for example, UreSil Ulti-flo Large Bore non-locking catheters (REF LBNL-series) to connect to a Gravity Drainage or Suction Drainage bag or system with standard luer. INSTRUCTIONS FOR USE: Used with funneled catheters, for example, UreSil Ulti-flo Large Bore non-locking catheters (REF LBNL-series), to connect to a Gravity Drainage or Suction Drainage bag or system with standard luer. CONTRAINDICATIONS: Currently no specific contraindications are known for this accessory for a medical device. WARNING: The reuse of this single-use device can lead to patient infection and/or device malfunction. Sterile if package is unopened and undamaged. Do not use if the sterile package is damaged or is unintentionally opened before use. SAFE DISPOSAL: Dispose of used device in container marked for biohazard (i.e. contaminated with potentially infectious substances of human origin). REPORTING SERIOUS INCIDENTS: Any serious incident that has occurred in relation to the device should be reported to UreSil as the manufacturer and the Competent Authority of the Member State in which the user and/or patient is established.