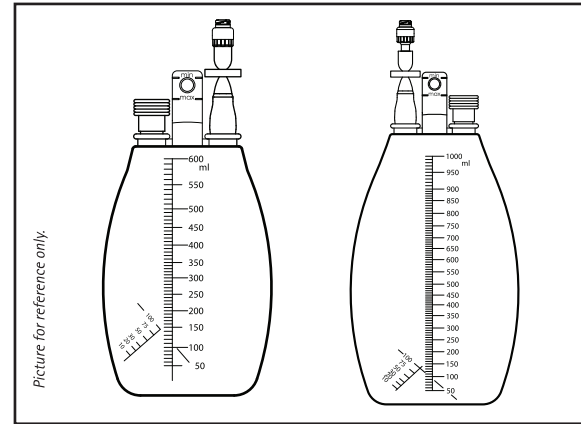


ASEPT[®] 600ml or 1,000ml Evacuated Drainage Bottle without Drain Line

INSTRUCTIONS FOR USE



WARNING: This product contains chemicals known to the State of California to cause cancer and birth defects or other reproductive harm.

Store at room temperature

This product contains dry natural rubber (latex)



Do not reuse



Do not resterilize



Do not use if package is damaged



Manufacturer



Keep away from sunlight



Consult instructions for use

REF

Catalog number

LOT

Batch code



Not made with DEHP

STERILE EO

Sterilized using ethylene oxide

Rx only



Keep dry



Use by

ASEPT[®] is a registered trademark of pfm medical, inc.

Interventional Systems
B | BRAUN

Distributed by:
B. Braun Interventional Systems Inc.
824 Twelfth Avenue
Bethlehem, PA 18018
www.bisusa.org



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Made in USA or Germany

Interventional Systems
B | BRAUN

Instructions for Use

ASEPT® 600ml or ASEPT® 1,000ml Evacuated Drainage Bottle without Drain Line

Contents of unopened, undamaged package are:

STERILE

Disposable - This device is intended for one use only.

Do not reuse or resterilize. Sterilized with ethylene oxide, sterile fluid path.

PRODUCT DESCRIPTION:

The ASEPT® 600ml or 1,000ml Evacuated Drainage Bottle without Drain Line is a vacuum based suction device for drainage purposes. The bottle comes with a needle-free valve attached to the bottle. The following instructions are for the ASEPT® 600ml or 1,000ml Evacuated Drainage Bottle.

The ASEPT® 600ml or 1,000ml Evacuated Drainage Bottle contains the following items:

One 600ml or 1,000ml vacuum bottle with the following features:

- Bottle scaled in 50ml increments
- Green vacuum indicator
- Vacuum release clamp
- Needle-free valve

GENERAL INFORMATION:

CAUTIONS:

- The green indicator should be down and level with the "max" line; this indicates that the inside of the bottle contains a vacuum
- When the green vacuum indicator reaches the position "min" a change of bottle is necessary
- Federal Law (USA) restricts this device to sale by or on the order of a physician

INDICATION:

The ASEPT® 600ml or 1,000ml Evacuated Drainage Bottle without Drain Line is a non-powered, single patient, portable suction device used to aspirate, remove, or sample body fluids.

APPLICATION:

Using the ASEPT® 600ml or 1,000ml Evacuated Drainage Bottle without Drain Line.

Follow these step-by-step directions.

1. Pick up bag that contains the ASEPT® 600ml or 1,000ml Evacuated Drainage Bottle without Drain Line. Look at the green vacuum indicator on the top of the bottle. This indicator tells you if there is a vacuum in the bottle. If the indicator is up and level with the "min" line then discard the bottle and use another.
2. Open the peel package under sterile conditions and take out the ASEPT® 600ml or 1,000ml Evacuated Drainage Bottle without Drain Line.
3. Swab the needle-free valve with a disinfectant and use a luer connector to connect to the needle-free valve. Twist all the way down ensuring the connection is secure.
4. Slide the white slide clamp on the green tubing until it no longer pinches the green tube closed. The fluid will flow into the ASEPT® Drainage Bottle with Drain Line.
5. After the drainage procedure; disconnect the drain line from the needle-free valve.
6. To sample fluid, swab the valve with disinfectant and use a needle-free syringe to access the valve. Hold the bottle upside down and remove the fluid.
7. Dispose of the used ASEPT® 600ml or 1,000ml Evacuated Drainage Bottle without Drain Line according to facility policies and procedures.

STERILITY:

This device has been sterilized, is for single use only, and is not to be reused.

As long as the packaging remains sealed and uncompromised the contents within each package are sterile. B. Braun Interventional Systems will not accept, exchange or credit any product that has been opened but not used by the patient or purchaser.

WARRANTY: B. BRAUN INTERVENTIONAL SYSTEMS INC. WARRANTS THAT THIS PRODUCT WAS MANUFACTURED ACCORDING TO APPLICABLE STANDARDS AND SPECIFICATIONS. PATIENT CONDITION, CLINICAL TREATMENT AND PRODUCT MAINTENANCE MAY AFFECT THE PERFORMANCE OF THIS PRODUCT. USE OF THIS PRODUCT SHOULD BE IN ACCORDANCE WITH THE INSTRUCTIONS PROVIDED AND AS DIRECTED BY THE PRESCRIBING PHYSICIAN.