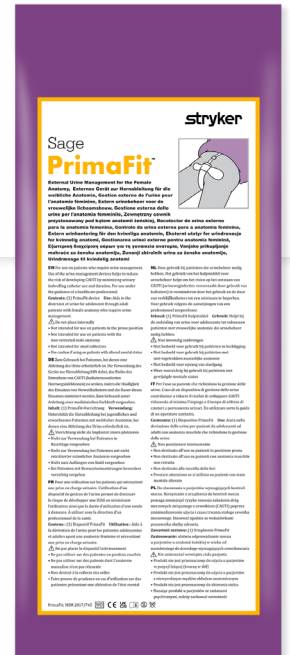


Sage

PrimaFit™

External Urine Management for the Female Anatomy



Prepping

- Perform skin assessment and document per hospital protocol.
- Clean, prep, and dry the patient's genital area from front to back per hospital protocol.

Applying

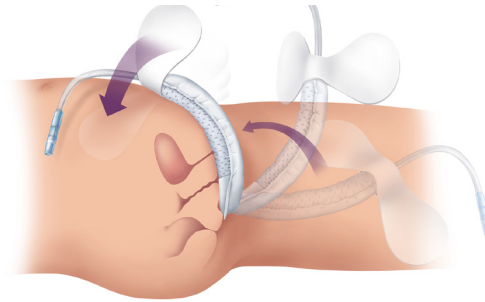
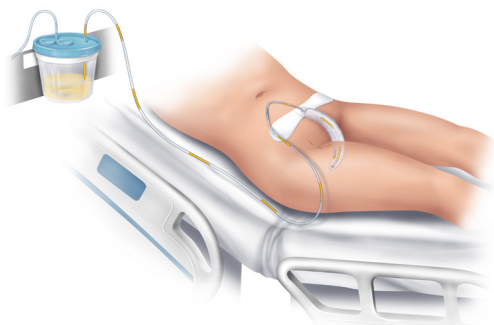
- Check patient positioning. Place patient in supine (flat) position when placing the device. After the device is placed, the head of bed can be raised.
- Align tapered end with perineum and place device between the labia and against the urethral opening.
- Make sure the fabric is covering the urethral opening.
- Use the date and time label to mark when device is placed on patient.

Maintaining

- Ensure suction is set on low, continuous suction at a **minimum of 8kPa, 80mmbar or 60mmHg.**
- Attach suction tubing to the device at the suction tubing adapter.
- Assess the device periodically to ensure proper placement, particularly after turning or repositioning the patient.
- Replace the device every 12-24 hours or if soiled with stool or bodily fluids other than urine.
- Remove the device before patient ambulation.

Placement guide

Flexible fit contours to patient anatomy and maintains shape for the duration of use



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A healthcare professional must always rely on his or her own professional clinical judgment when deciding whether to use a particular product when treating a particular patient. Stryker does not dispense medical advice and recommends that healthcare professionals be trained in the use of any particular product before using it.

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PrimaFit is a Class I medical device CE.

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