AN OSTOMY is a surgically created diversion for feces or urine after removal of part or all of the intestine or bladder. According to the United Ostomy Associations of America (UOAA), an estimated 750,000 Americans are living with an ostomy and 130,000 new ostomy surgeries occur in the United States annually.

Due to the complex nature and labor intensity of ostomy management, many hospitals employ certified wound, ostomy, and continence nurses (CWOCN) as resource persons. Also known as WOC nurses, they have advanced education in these specialties. But even if your hospital has a nurse ostomy specialist (and especially if it doesn’t), you need to be able to care for patients who have an ostomy. This article will provide the nuts and bolts so you can do just that.

**About ostomies**
Whether temporary or permanent, an ostomy may be created for such indications as cancer, inflammatory bowel disease (Crohn’s disease or ulcerative colitis), trauma, bowel obstruction, and bowel perforation. Common types of diversions include ileostomy, colostomy, and ileal conduit. (See *Understanding ileostomies, colostomies, and ileal conduits*.) The diversion may be incontinent or continent; this article discusses incontinent diversions only.

**Preoperative preparation**
Before surgery, the nurse ostomy specialist or another nurse meets with the patient and home caregiver to provide information and support and to assess for an optimal surgical site for stoma creation. This critical session also provides an opportunity for patients to express their thoughts and feelings about the upcoming surgery and ostomy.

**Postoperative care**
After surgery, you’ll need to identify which type of ostomy the patient has, its indication, and whether it’s temporary or permanent. This information helps you gauge the patient’s overall health status, guide interventions, and determine educational needs. If necessary, you may need to check another source, such as the operative notes, to get answers to these questions. (Note: You can’t determine the type of os-
tomography a patient has based just on its abdominal location.

You also need to understand various ostomy appliances and accessories, how to assess the patient, and principles of emptying and changing the pouch.

Ostomy appliances

The nurse ostomy specialist chooses the initial appliance (pouching system) after measuring the stoma. (Measurement devices are available from appliance companies.) At each appliance change, you should measure the stoma because swelling may change its size or position. Swelling typically subsides over the first 6 weeks after surgery.

Wear time may vary; in most cases, it’s 5 to 7 days for a colostomy and 3 to 4 days for an ileostomy or ileal conduit. After the first 6 weeks, the patient may change to a different appliance with professional guidance.

The ostomy appliance consists of a skin barrier and pouch to contain fecal or urine effluent. (See Types of skin barriers.) Disposable and waterproof, the appliance is designed to contain odor and has either a drainable clamp or Velcro closure or is a nondrainable closed-ended pouch.

Some appliances are clear; others are opaque. An opaque appliance may make it difficult to center the pouch directly over the stoma. All pouching systems are equally effective, so the choice depends largely on patient preference.

The ostomy appliance may have one or two pieces.

- A one-piece system has a single seal, with the skin barrier permanently attached to the pouch. Appropriate for patients with a flat peristomal profile, it entails one less application step than a two-piece system.
- With a two-piece system, the pouch can be detached from the skin barrier and discarded.

Ileostomies and colostomies are fecal diversions. Created from the last section of the small bowel (ileum), an ileostomy is typically created in the right lower abdominal quadrant. Once it begins to function (usually in 1 to 3 days), effluent (stool) volume may be high, but in most cases will subside in a few weeks. At first, effluent volume is about 1,200 mL/24 hours, and then it decreases to around 800 to 1,000 mL/24 hours. Although the stoma is created from the ileum and output initially is liquid, the body learns to compensate and eventually stool consistency may become more like toothpaste.

A colostomy is created from any part of the large bowel—ascending, transverse, descending, or sigmoid colon. Expected stool volume is approximately 600 to 1,000 mL/24 hours. The closer the stoma to the end-point of digestion (descending or sigmoid colon), the more formed the stool is.

- With a colostomy at the ascending colon, stool consistency is similar to that created by an ileostomy.
- A colostomy at the transverse colon leads to a stool consistency like toothpaste.
- A colostomy at the descending colon causes semiformal stool.

An ileal conduit is a urinary diversion. If the bladder is removed, a cystectomy is done, followed by a 12- to 18-cm (5- to 7-inch) resection of the ileum, in which a conduit is formed from the ureters to the outer abdomen. The proximal end of the ileal conduit is closed and the distal end serves as the stomal opening on the outside of the abdomen; the ureters are anastomosed to the ileal conduit segment. Urinary volume should be at least 800 mL/24 hours.
instead of emptied. However, it requires an extra step—re-attaching the pouch to the skin barrier.

**Ostomy accessories**

Various ostomy accessories promote comfort, adherence, and healthy peristomal skin. Skin-barrier paste and powder, as well as barrier rings, wafers, and strips, are made from adhesive pectin. Skin-barrier prep and powder are available as sprays, wipes, or “lollipop” sticks. Applied directly to skin around the stoma before barrier or pouch application, they protect the skin by preventing effluent from leaking and touching the skin under the appliance. (Caustic stool enzymes in effluent can harm peristomal skin.)

Clinicians must read all manufacturer directions before using these products. If peristomal skin is denuded (a condition called epidermal skin stripping), powder can be sprinkled directly on the denuded skin and sealed by dabbing at it with the corner of skin-barrier film prep—a technique called crusting.

Other ostomy accessories include:

- ostomy belts, which help secure the appliance
- binders, which support a peristomal hernia
- wraps, which can provide a flatter body contour
- special swimwear and intimate apparel that allows for privacy.

**Patient assessment**

When assessing the patient with a new ostomy:

- inspect the appliance to verify that it’s secure and functioning properly
- check for effluent leakage
- note the date of the last pouch change
- evaluate output character and volume
- check for flatus.

When emptying and changing the pouch, inspect the stoma, noting its anatomic location and proximity to drains or the surgical incision, as well as stoma characteristics and peristomal skin condition. Normally, the stoma is red, moist, and firm and protrudes 2.5 cm (1 inch) above the skin surface. The opening should point directly upward.

Stoma shape and size depend on how the stoma was created and its location in the GI tract. Stomal discoloration, inadequate output, absent or diminished bowel sounds, and abdominal distention or tenderness may indicate stoma dysfunction, poor perfusion, or both. If these signs or symptoms occur, a surgical consult may be necessary. Be aware that slight bleeding may occur with gentle cleansing because stomal mucosal tissue has a rich blood vessel supply.

**Emptying and changing the pouch**

The pouch should be emptied when it’s one-third to one-half full. In a high-volume system, such as a new ileostomy or ileal conduit, an appliance with an output spout may be connected by a conduit connection to a urinary leg-bag drainage system, allowing for a larger reservoir and less frequent changes.

Ideally, patients should empty the pouch while standing at or sitting on a toilet with their legs open. Patients with impaired mobility may lie on their side and drain the pouch into a urinal or wash basin, then dispose of the output in a toilet.

Basic pouches should be changed at least weekly or whenever leakage occurs. High-volume output systems, such as ileostomies and ileal conduits, need to be changed more often. (See Changing the ostomy appliance.)

**Educating patients and home caregivers**

The initial phase of a new ostomy can be overwhelming for patients and caregivers. To build their confidence, work closely with them, conveying confidence and patience. Teaching points to cover include ostomy type, expected output, the need to alert nursing staff of leakage and other issues. (continued on page 10)
Changing the ostomy appliance

Follow these steps when changing the patient’s ostomy appliance.

1. Gather all needed supplies.
2. Perform hand hygiene.
3. Assemble and open supplies on a clean surface near the patient.
4. Stretch back the skin-barrier opening, or cut an opening on a starter hole and peel back to remove. Connect the skin barrier to the pouch.
5. Tuck an incontinence pad inside the patient’s clothing to prevent soiling.
6. Set up a mirror on the bedside table so the patient can observe the procedure.
7. Don gloves.
8. Remove the old appliance by holding the skin taut with one hand and gently pulling back the skin with your other hand (push-pull technique).
9. Discard the old pouching system.
10. Clean the stomal and peristomal areas with moistened paper towels. Avoid using soap or baby wipes, as they may leave a residue that can interfere with the appliance seal.
11. Have gauze ready when preparing the peristomal area to wick leaking effluent.
12. Inspect the stoma and peristomal areas for abnormalities and irritation.
13. Measure the stoma with a measuring guide to determine if the patient needs a larger or smaller appliance.
14. If you note peristomal skin irritation, perform the crusting technique, using skin-barrier powder and prep.
15. Apply skin-barrier prep to remaining peristomal skin and let dry for 2 to 3 minutes.
16. If using a barrier ring, apply so it fits directly around stoma.
17. With a two-piece appliance, attach the skin barrier to the pouch at the flange (where the barrier and pouch connect) by snapping it in place. Then check for securement, if you didn’t do this earlier.
18. Remove adhesive backing from the barrier and center the barrier or pouch directly over the stoma. Secure adhesive to skin around the barrier.
19. Have the patient put one hand, palm side down, directly over the pouch adhesive for 5 to 10 minutes to warm the materials, allowing for a stronger seal.
20. Close and secure the bottom opening of the pouch.
21. Optional: Frame the appliance with tape, or apply an ostomy belt.
22. Remove gloves and perform hand hygiene.

From anxiety to empowerment

Patients with new ostomies face an initial period of challenging psychological and social adjustments. Major concerns include altered self-image, fear of rejection, intimacy changes, and fear that the ostomy appliance will lose adherence in a public or social situation.

Provide emotional support and urge patients to express feelings of age or seal compromise, and the importance of emptying the pouch when it’s one-third to one-half full. Also teach patients and caregivers about an appropriate diet to follow. (See Dietary considerations.)

Determine the patient’s readiness to learn about the ostomy. For patients who won’t look at or touch the pouch, use a gradual, supportive, step-by-step approach. For those who seem receptive to seeing, touching, and learning about the pouch, explain the role of each ostomy item and urge the patient to touch and practice with these items. (Manufacturer-supplied ostomy models can aid this process.) Once the patient is comfortable with the pouch, demonstrate a pouch change; if the patient expresses readiness, he or she may participate.

At the next pouch change, encourage the patient to perform a return demonstration with coaching. Have the patient open and close a sample pouch, attach a pouch to a skin barrier, and practice emptying his or her own pouch. Reassure patients that their appliance will adhere successfully even if challenges occur at first.

Many patients and caregivers are far less anxious after the return demonstration. At subsequent pouch changes, supervise or help the patient or caregiver, as needed.

At discharge, the patient should be able to empty the pouch independently. Otherwise, explore alternatives, such as a family member or caregiver, who can help the patient. Provide written materials, resource information, and peer-support referral, all available from the UOAA (ostomy.org), Wound Ostomy and Continence Nurses Society (wocn.org), and Ostomy Secrets (ostomysecrets.com). As appropriate, arrange for a home health nursing referral for additional support. Other helpful resources include ostomy clinics and certified ostomy nurses who practice in the community; for information on the latter, contact the WOCN Society.
anxiety and fear. Teach them how to take control of their situation to improve their quality of life. With emotional support, information, and practice, ostomy patients can feel empowered and achieve positive outcomes.

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Selected references
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Dietary considerations
Most patients can resume a normal diet after ostomy surgery but should limit their intake to small, frequent meals with low fiber intake for the first 2 weeks (due to bowel edema). Encourage them to chew foods thoroughly to maximize healthy digestion.

During the initial high-volume effluent period, an ileostomy must be monitored closely because of the risk of fluid and electrolyte imbalances. Patients with loose stools may benefit from stool-thickening foods, such as applesauce, bananas, rice, peanut butter, and bread. They should chew these foods, particularly those high in fiber, thoroughly to avoid bowel blockage. Patients with ileal conduits don’t require dietary modification but should drink plenty of fluids, especially water, to help prevent mucus buildup in the urine.

Know that certain foods and beverages, including dairy foods, onions, cabbage, broccoli, mushrooms, beans, cucumbers, and alcohol, may trigger gas. Gas causes the pouch to expand; to release gas, instruct the patient to open the pouch at the flange (where the pouch and skin barrier connect). To help reduce gas, advise patients to avoid chewing gum, drinking carbonated beverages, and using beverage straws.

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