By 2050, an estimated 27 million people will need some type of long-term care. Home healthcare and hospice agencies are the major providers of community-based long-term care. Currently, about 7.6 million people receive community-based care for post-acute and chronic conditions, often with multiple co-morbidities. This number is expected to increase as the population ages. This issue of Perspectives focuses on best practices in the management of home care patients, in particular patients with either an indwelling catheter or a trachecotomy.

Although there is ample research-based evidence regarding indwelling urinary catheter management in acute and long-term care settings, there is limited home care information. Dr. Wilde’s and Mr. Zhang’s article describes the application of evidence-based practices for the home care patient.

Care of the trachecotomy in the home is a growing trend due to the increased efforts to transition patients to less costly points of care, along with the technologic advances that allow caregivers to deliver limited forms of medical care in the home.

### Advisory Board

**Jan Foster**, RN, PhD, MSN, CCRN  
Associate Professor of Nursing  
Texas Woman’s University, Houston, TX

**Mikel Gray**, PhD, CUNP, CCN, FAAN  
Professor and Nurse Practitioner  
University of Virginia  
Department of Urology and School of Nursing, Charlottesville, VA

**Tim Op’t Holt**, EdD, RRT, AEC, FAARC  
Professor, Dept. of Respiratory Care and Cardiopulmonary Sciences  
University of South Alabama, Mobile, AL

**Paul K. Merrel**, RN, MSN, APN-2  
Advance Practice Nurse, Adult Critical Care  
University of Virginia Health System, Charlottesville, VA

**Jennifer A. Wooley**, MS, RD, CNSD  
Clinical Nutrition Manager  
University of Michigan Health System, Ann Arbor, MI

Dale Medical is committed to nursing education and is the proud sponsor of Perspectives, a source of free, quality CEs for nurses for the past 15 years.

---

**Best Practices in Managing the Indwelling Urinary Catheter for the Homecare Patient**

By Mary H. Wilde, RN, PhD and Feng Zhang, RN, BS

Indwelling urinary catheters can be used for short periods of time, such as post-hospitalization, or indefinitely for persons with chronic urinary retention. Long-term catheters (LTC) are reserved for people who cannot perform intermittent catheterization, have no one to do it for them, or cannot manage any other way. Also, in certain individuals a catheter might be a quality of life decision making it easier to travel or work. Nevertheless, use of an indwelling urinary catheter puts the person at risk for persistent catheter-related problems. The purpose of this article is to address the nursing management of people with indwelling urinary catheters, primarily those with long-term catheters, and to suggest best practices based on evidence to minimize or prevent catheter related problems.

### Short and Long-term Catheters Defined

Short-term catheters are defined as catheters that are used for less than one month. In actual home care practice, this time frame could be extended to a few months before the catheter is discontinued. Sometimes a “trial without a catheter” by removal and monitoring voiding and residual urine is not successful and the catheter is reinserted. Short-term catheter users need to keep a closed system, which can minimize or delay bacteria in the urine (bacteriuria) and possibly symptomatic catheter-associated urinary tract infection (CAUTI); however all with a catheter will have bacteriuria after 30 days. Irrigations should not be done unless they are prescribed by a physician for a specific purpose such as removal of blood clots. The catheter type can include polytetrafluoroethylene (PTFE or Teflon)-coated latex, which is soft but provides some protection from latex. It can be used for up to a month. Coatings on catheters, such as silver alloy—but not silver oxide—might be of value in decreasing bacteria in urine for short-term use up to 2 weeks, but neither silver alloy nor antimicrobial catheters have prevented symptomatic CAUTI.

Long-term catheter use requires different considerations. First, the decision should be made about whether the catheter will remain in situ, and whether other options would be better. Persons with persistent retention, hand dexterity and mental capacity can be taught to use intermittent catheters, as well as family or caregivers. For men with severe incontinence and no retention, an external catheter could be an option.

Continued on page 7
Tracheostomy Care in the Home Care Patient

Tim Op’t Holt, EdD, RRT, AE-C, FAARC
Jennifer McDaniel, RRT-NPS
Kathleen Deakins, MSHA, RRT-NPS, FAARC

Tracheostomy, the creation of an opening in the neck into the trachea, is performed in the event of airway obstruction, prolonged endotracheal intubation, and for tracheobronchial hygiene. In adults, tracheostomy is done by traditional surgical opening of the trachea in the operating room and by the more recent percutaneous dilational tracheostomy technique at the bedside, usually in the intensive care unit. At a local hospital in south Alabama, it was reported that a total of 191 tracheotomies had been performed in 2010 and 2011. Of those, 79 (41%) patients were discharged with the tube in place. It may be common for a patient to be discharged with the tube in place, putting the burden for care on the family and the responsibility for preparing the family for this task on hospital educators. Care of the tracheostomy in the home is a growing trend due to increased economic pressures on acute-care medical facilities, efforts to transition patients to less costly points of care, technologic advances that allow caregivers to deliver limited forms of medical care in the home, and the growing social acceptance of persons with disabilities. When the tracheostomy is no longer indicated, the tube is removed (decannulation), and the patient resumes ventilation through the native airway.

Information from a large database of pediatric patients revealed that in 2009, 4341 tracheotomies were performed on children in the 46 states participating in the Healthcare Cost and Utilization Project (HCUP) Kid’s Inpatient Database, sponsored by the Agency for Healthcare Research and Quality. While data on the number of children discharged with a tracheostomy are unavailable, many children are discharged to home with a tracheostomy, as evidenced by the numbers of health care workers engaged in teaching caregivers how to care for children with tracheostomy in the home.

Indications for Tracheostomy
Tracheostomy is indicated for upper airway obstruction, prolonged mechanical ventilation, and neuromuscular disease. Airway obstruction may be caused by or related to infection, trauma, tumor, foreign body, obstructive sleep apnea, or tracheal stenosis. Once the upper airway obstruction is bypassed by the tracheostomy, the tracheostomy tube is inserted and the patient may be able to ventilate spontaneously. After a period of time (2 days to several weeks of oral intubation) a tracheostomy is considered for patients requiring prolonged mechanical ventilation, the timing of which is a topic of considerable controversy. In patients with respiratory failure, the tracheostomy facilitates tracheobronchial hygiene and may speed the ventilator liberation process, since airway patency is better assured. Likewise in neuromuscular disease, a tracheostomy facilitates airway access for hygiene and attachment of oxygen and ventilatory devices. The long-term presence of a tracheostomy is dictated by the failure of an acute tracheostomy situation to resolve. For example, a post-trauma comatose patient may not regain consciousness and therefore has a chronically compromised airway. Attempts to decannulate such a patient often results in the need for re cannulation.

In premature infants, tracheostomy has been reported in approximately 2.8% of very low birthweight infants (361-1500 g), or about 1500 children/year. This is to facilitate long-term mechanical ventilation, often associated with bronchopulmonary dysplasia. Tracheostomy is also indicated in infants born with severe maxillofacial abnormalities or neurologic deficiency. Most children requiring a tracheostomy are younger than 1 year of age. In these children, tracheostomy is associated with better growth and development, better development of facial expression and phonation, and improved ability to learn sucking and swallowing. They are also able to be cared for in the home environment as opposed to patients who remain orally intubated. Other indications include trauma (usually closed head injury and need for mechanical ventilation), postoperative complications following major cardiothoracic surgery, and infection/sepsis-related disease.

Issues in the Care of a Patient with a Long-Term Tracheostomy
There are several issues in the care of the patient with a long-term tracheostomy. These issues are central to the airway itself and those pertaining to the mechanical ventilation that often accompany the presence of the tracheostomy tube. Mechanical ventilation issues are beyond the scope of this presentation. However, when considering the tracheostomy alone, cuff pressure, cuff leak, tube dislodgement, tube occlusion, infection, changing the tube, suctioning, and promoting communication are important issues for the patient and caregiver.

Cuff pressure and cuff leak are...
In patients who are not being ventilated, the cuff may be left deflated if the patient has good upper airway of this issue is beyond the scope of this paper. Speech is facilitated by partial deflation of the cuff during mechanical ventilation. In patients who are not being ventilated, a fenestrated tracheostomy tube can facilitate speech during exhalation. A speaking valve, such as the Passy-Muir valve allows airflow through the tube during inspiration, then routes the air through the vocal cords during expiration. A referenced, full explanation of these issues can be found elsewhere.4

Patient and Caregiver Education for Children with a Tracheostomy, from the Therapist’s Perspective

Note: Please refer to the “Parent Guide to Trach Care” at the following website: www.perspectivesinnursing.org. Most of these comments on care for the child with a tracheostomy also apply to care of the adult with a tracheostomy.

The teaching of home tracheostomy care should ideally begin prior to the procedure. Consideration should be given to the family’s level of education and their feelings of apprehension or anxiety regarding the impending procedure. Talking with the family about “what is a tracheostomy” and assuring them of their ability to care for their child is extremely important. Answer their questions truthfully and honestly and in the simplest manner possible. Be positive in all interactions with the family. Try not to overwhelm the family at this time with too much information—there will be plenty of time after the surgery to emphasize skills.

Some parents/caregivers wish to see another child with a tracheostomy. If there is another child in the hospital with a tracheostomy, you must get the parent’s permission before bringing your new parents into the room. Also, be especially cautious with what type of patient you present to the parents. A parent with a child with an acute upper airway obstruction who is fairly normal in other aspects might become distressed at seeing a child in a chronic vegetative state with a continuously bubbling tracheostomy.

Leave educational booklets and information and encourage the parents to look over and write their questions down prior to watching the tracheostomy care video. It is very important during this anxious period for the therapist to be available to answer questions and offer reassurances. Encourage the review of all training materials. If parents have access to a computer, make them aware of Aaron’s Tracheostomy Page at www.tracheostomy.com.

During the first 24 hours after the surgery, parents may or may not be able to hold the child (usually not if on ventilator). Encourage parents to observe care during this time, but do not force this issue. Remember, to the parents the tracheostomy looks like a big, gaping, oozing hole through which a plastic tube has been inserted. Give the parents time to adjust and accept the tracheostomy.

During the first week postoperatively, encourage parents to observe and participate in tracheostomy care/tube changes/suctioning/manual ventilation. Remember it is a scary thought to have to insert something inside your child, especially if it makes the child cry while you are doing it.

Once the surgeon has changed the tracheostomy tube after at least 1 week, training for parents and caregivers should be increased. Meet with the social worker at this time to coordinate training schedules, prepare an equipment list for home care companies, arrange rooming-in requirements and target a discharge date. Ideally, we would like to be able to discharge the patient within 1 week of the surgeon’s initial tracheostomy tube change or 2 weeks from the placement of the tracheostomy if that is the only apparent problem for the patient.

One of the first things to teach the parent is tracheostomy care and
Parents/caregivers should be taught how to assess their child’s respiratory status and what signs or symptoms may indicate respiratory distress. If the child is experiencing signs of respiratory distress, the child may need to be ventilated and suctioned, the tube may need to be changed, a breathing treatment may be indicated, or the child may need to be transported to the hospital. Once parents have changed the tracheostomy tube at least once and can state or recognize signs of respiratory distress, begin teaching what to do in an emergency if the tube were to become plugged. Emergency tracheostomy tube procedures are probably the most important training the parents will receive. Review them daily with the parents/caregivers until the child is discharged. Caregivers need to be able to tell you what to do and do it without hesitation.

Once parents are comfortable with emergency tracheostomy procedures, you may begin teaching them CPR using a mannequin for practice. The hospital educator is responsible for teaching and checking off CPR skills. Always make them do the 4 steps in emergency tracheostomy procedures as their initials steps of CPR. It does absolutely no good to do compression if the airway is not patent.

Review the contents of the travel bag with parents and emphasize the importance of taking the bag everywhere the child goes. This should be a completely different bag than the diaper bag. Parents should be able to state the contents of the bag.

If the child is old enough to begin making sounds (usually 6-9 months), they should be considered for a speaking valve. Consult with ENT and speech therapy before placing the valve. (A physician order is necessary to use a valve.) Because it is a 1-way valve with the patient breathing in through the tracheostomy and out through the mouth there must be a leak present around the tracheostomy tube for air movement. Ideally we would like to place the valve in the hospital before the patient is discharged, but the patient can be brought in as an outpatient for valve placement.

Parents should be encouraged to continue working on the above technical and decision-making skills. Coordinate with social services a discharge planning meeting to occur during this time to involve parents and all disciplines (social services, respiratory therapy, physical and occupational therapy, speech therapy, nutritionist, physicians and home care coordinators, nursing and durable medical equipment [DME] providers). Have a final equipment list prepared for the DME provider. Finalize rooming-in requirements (coordinate with the DME for delivery of their equipment and of their training needs) and set a discharge date.

Parents should be now ready to change out the tracheostomy tube. It is probably best to let them change the tracheostomy tube with your assistance. Teach the parents to gather all the equipment and prepare the new tracheostomy tube. Once equipment has been gathered and the tracheostomy tube change has been performed, the child should be monitored for color and/or respiratory distress. If there is ever any question about where the tube is, ventilate the patient and look for chest movement. If in doubt, remove the tube and replace it. Once the tracheostomy is in, the tracheostomy tube ties are fastened, the tube is suctioned as needed and the tracheostomy site cleaned. The primary caregiver should change the tracheostomy tube 3 times before discharge and the secondary caregiver should do the same at least once.

Parents/caregivers should continue working on the above technical and decision-making skills. Coordinate with social services a discharge planning meeting to occur during this time to involve parents and all disciplines (social services, respiratory therapy, physical and occupational therapy, speech therapy, nutritionist, physicians and home care coordinators, nursing and durable medical equipment [DME] providers). Have a final equipment list prepared for the DME provider. Finalize rooming-in requirements (coordinate with the DME for delivery of their equipment and of their training needs) and set a discharge date.

Parents should be now ready to change out the tracheostomy tube. It is probably best to let them change the tracheostomy tube with your assistance. Teach the parents to gather all the equipment and prepare the new tracheostomy tube. Once equipment has been gathered and the tracheostomy tube change has been performed, the child should be monitored for color and/or respiratory distress. If there is ever any question about where the tube is, ventilate the patient and look for chest movement. If in doubt, remove the tube and replace it. Once the tracheostomy is in, the tracheostomy tube ties are fastened, the tube is suctioned as needed and the tracheostomy site cleaned. The primary caregiver should change the tracheostomy tube 3 times before discharge and the secondary caregiver should do the same at least once.

Parents/caregivers should be taught how to assess their child’s respiratory status and what signs or symptoms may indicate respiratory distress. If the child is experiencing signs of respiratory distress, the child may need to be ventilated and suctioned, the tube may need to be changed, a breathing treatment may be indicated, or the child may need to be transported to the hospital. Once parents have changed the tracheostomy tube at least once and can state or recognize signs of respiratory distress, begin teaching what to do in an emergency if the tube were to become plugged. Emergency tracheostomy tube procedures are probably the most important training the parents will receive. Review them daily with the parents/caregivers until the child is discharged. Caregivers need to be able to tell you what to do and do it without hesitation.

Once parents are comfortable with emergency tracheostomy procedures, you may begin teaching them CPR using a mannequin for practice. The hospital educator is responsible for teaching and checking off CPR skills. Always make them do the 4 steps in emergency tracheostomy procedures as their initials steps of CPR. It does absolutely no good to do compression if the airway is not patent.

Review the contents of the travel bag with parents and emphasize the importance of taking the bag everywhere the child goes. This should be a completely different bag than the diaper bag. Parents should be able to state the contents of the bag.

If the child is old enough to begin making sounds (usually 6-9 months), they should be considered for a speaking valve. Consult with ENT and speech therapy before placing the valve. (A physician order is necessary to use a valve.) Because it is a 1-way valve with the patient breathing in through the tracheostomy and out through the mouth there must be a leak present around the tracheostomy tube for air movement. Ideally we would like to place the valve in the hospital before the patient is discharged, but the patient can be brought in as an outpatient for valve placement.

Parents should be encouraged to treat the child as normally as possible. Parents may obtain hints for daily living from the reading materials, talking with other parents or children with a tracheostomy, or going online to Aar-
on’s Tracheostomy Page, cited above.

Once you have insured that all training has been completed and documented, it is now time to send the baby home. Answer any questions the parent/caregivers may have or address any problems that may have arisen during rooming in. Contact the DME home therapist to make sure all equipment is in place and nothing else is needed. Work with the social worker, resident and nursing to write prescriptions for equipment and therapy. Review home nursing orders to ensure respiratory concerns are addressed. Congratulate the caregivers, wish them well and walk the patient and family to the door.

**Securement and Decannulation of the Tracheostomy Tube**

Three materials are used for securing the tracheostomy tube to the patient: cotton twill tape, commercially available Velcro® and felt (fabric) tracheostomy securement devices, and a beaded chain. An American Thoracic Society committee looking into the use of these three devices in children made the following observations.\(^6\)

Cotton twill tape may trap moisture, becomes soiled frequently, may unravel, and subsequently irritate the skin. Velcro devices have less tendency to abrade the skin, as does twill tape. They are convenient and easily adjusted. One-person tie changes are easier with Velcro ties than with other tie materials (Figure 1). They are wider than cotton twill and chain, so they have less tendency to abrade the skin. They are easily adjusted and because of their padding material, should be quite comfortable. There is third-party coverage for these securement devices. A stainless steel beaded chain is durable, does not trap moisture, and maintains a constant tension. Any securement device is kept tight enough to allow slipping a finger between the neck and device material. The consensus was that the most important aspect of choosing a tracheostomy tie is how well the tie can be secured, rather than the material it is made from.\(^5\)

While there does not seem to be any similar data available for tracheostomy securement in adults, these same principles should apply.

The removal of the tracheostomy tube is referred to as decannulation. This implies that it is the intention to not reinsert the tube, as is the case with a tube change. Because of equipment concerns and the potential for a need for emergency airway management, decannulation is usually not done at home.\(^6\) None of the recommendations home decannulation.

**Special Tracheostomy Issues for Children**

Like adults, children sometimes require placement of a tracheostomy when there is a need for a short term or permanent airway, or if there is a need for long-term invasive ventilation. The indications for tracheostomy in children differ from adult patients and are widespread. Of the 16 characteristic indications for pediatric tracheostomy, the most common are airway obstruction, need for airway protection due to aspiration or inability to manage secretions, evidence of subglottic stenosis, chronic lung disease patients requiring long term mechanical ventilation, and the presence of congenital abnormalities.\(^8\)

Unlike adult patients who may be considered candidates for tracheostomy after approximately 1 week of invasive mechanical ventilation, children are often not considered candidates for tracheostomy until much later in their course, with the exception of those with anatomic airway abnormalities. Davis indicated that the average time of tracheostomy insertion in children was at about 65 days following initial intubation; and is often longer in some cases.\(^8\) Despite the variation in length of time a tracheostomy is needed, about two-thirds of pediatric patients are eventually decannulated.\(^8\) Because of the unpredictable duration of pediatric tracheostomy, patients and families/caregivers begin preparations for transition to home as soon as feasibly possible.

Children often remain in the intensive or specialized care units until the first tracheostomy change is completed: around 5-7 days after insertion.\(^1\) Close observation is required because a large number of pediatric tracheostomy tubes are small and uncuffed, and there is a greater chance for dislodgement or decannulation. It is recommended that specialized trained caregivers such as respiratory therapists or nurses trained in tracheostomy care assist the surgeon during the initial tracheostomy tube change, as described in the previous section.\(^1\) Patients are typically transitioned to a lower level of care and an organized approach to patient and caregiver training begins.

Hospital stays for children with a tracheostomy vary amongst institutions. Some programs report length of stay at 5 to 472 hospital days while others are 14 to 280 days from initial admission.\(^6\)\(^8\) Causes for these variations are multifactorial. Proactively preparing the tracheostomy patient and family for discharge and care in the home setting is highly recommended.\(^1\) Successful discharge planning programs have had positive outcomes if education initiatives were started before the initial surgery.\(^1\) Because patient’s medical issues, or family social concerns often create barriers and play a role in delaying discharge, attempts are made to keep caregivers on an education and training timeline. This allows time for home assessment, equipment selection, repetition of skills and retention of concepts presented.\(^8\) The need for nursing care in the home frequently contributes to a prolonged

---

**Figure 1. Dale Tracheotomy Tube Holder (Pediatric)**

**Figure 1a. Dale Tracheotomy Tube Holder**
hospital stay. Pediatric patients requiring home nursing care are on mechanical ventilators, have an unstable airway or have multiple complex medical needs in addition to tracheostomy care. In some cases, an average of 8-24 hours of nursing care may be granted based on these requirements. The level of nursing care varies based on degree of medical complexity, medical insurance benefits or state-based resources. Orchestrating these services is the greatest challenge and may consume a significant amount of time in this phase. In addition, complex training requirements and patient and family education require additional resources. Most importantly, coordinated care, streamlined education and training and shortened the length of stay from tracheostomy insertion to discharge are primary objectives of these programs.

In 2000, discharge planning programs for pediatric patients were initially created based on prior recommendations from the American Thoracic Society’s Care of the Child with Chronic Tracheostomy. Since then, much attention has been focused on standardizing care of the pediatric tracheostomy patient prior to discharge. Graf et al attempted to emulate this by implementing a family education program for pediatric tracheostomy patients. The Texas Children’s Hospital program reduced the length of stay following tracheostomy to a mean of 6.5 days following an educational program (5-114 days total length of stay). In 2012, the American Academy of Otolaryngology-Head and Neck Surgery created a clinical consensus statement to provide updated recommendations for pediatric and adult patients with a tracheostomy. Some recommendations have been tested and have since become expectations of care for these patients. To prepare families or caregivers for caring for the pediatric tracheostomy patient at home, 2 caregivers are identified as soon as the decision is made to proceed with the procedure. Basic education is disseminated to caregivers via handouts, such as a specialized tracheostomy care booklets (or electronic versions if desired) as described above. Upon gradual completion of these components, the caregiver is tested to assess knowledge and competency. The most unique element of a pediatric discharge planning program is a requirement for families to “room in,” or spend a 24-hour or other designated period of time taking care of the child independently as if they were in a home setting while having the ability to raise any final questions and calm their fears or uncertainties prior to discharge.

Pediatric tracheostomy patients are cared for by families and nurses in the home and monitored on a regular basis by respiratory therapists. Physicians assess the airway to determine if changes in airway size is necessary, make modifications in mechanical ventilator settings or verify the time for liberation from all components. If decannulation is being considered, the patient must be free of aspiration episodes, not require mechanical ventilator support, have completed a tracheostomy capping trial if appropriate, and have been evaluated by bronchoscopy within a few months of considering decannulation. Pediatric tracheostomy-ventilator clinics provide a vehicle to assess changes in the patient’s condition and allows for further evaluation. Streamlining pediatric tracheostomy care is an ongoing challenge and a unique opportunity to make a difference!

References

Timothy B. Op’t Holt, EdD, RRT, AE-C, FAARC, is Director of “Breath of Life” COPD and the Asthma Education and Therapy Program at Victory Health Partners Clinic in Mobile, Alabama. At the University of South Alabama, he is Professor, Department of Respiratory Care and Cardiopulmonary Sciences. He is the author or co-author of 8 books and 30 studies in journals and has presented over 35 papers at international conferences.

Kathleen M. Deakins, MSHA, RRT-NPS, FAARC is Clinical Manager, Women’s and Children’s Respiratory Care, Pediatric Pulmonary Function and Infant Monitoring Rainbow Babies & Children’s Hospital of University Hospitals, Cleveland, Ohio. She is an accredited neonatal-pediatric specialist with an interest in all facets of pediatric and neonatal respiratory care. She has published more than 50 publications on respiratory medicine and lectured widely in her field. She is also a member of several professional organizations dedicated to respiratory medicine. Ms Deakins lives in Chardon, Ohio.

Jennifer D. McDaniel, RRT-NPS is currently a respiratory therapist at the USA and Children’s Hospital In Mobile, AL. In addition to her clinical duties in the neonatal intensive care, Ms. McDaniel teaches neonatal and pediatric respiratory care. Ms. McDanels received her Bachelor of Science in Cardiorespiratory Care from the University of South Alabama.
they are commonly used in select patients in the United Kingdom (UK) and Europe. When a catheter is expected to be used indefinitely, it can be with the person for many years. In one study of 202 long-term catheter users, the mean duration of use was 6 years (SD 7) and the median was 3.25 years. Therefore, patients and their families need to learn how to manage the catheter between nurse visits.

**Catheter Selection**

For long-term catheter patients, the catheter type should be hydrophilic or coated with silicone. Pure silicone is not as pliable but it has a larger inner lumen which is of value for people with large amounts of sediment that blocks and causes frequent catheter changes. However, in one study, half the water evaporated within 3 weeks and 85% within 8 weeks. This can cause the catheter to “slip down,” causing irritation to the opening of the bladder and/or leaking (by-passing of urine). Latex allergy is of concern to many with long-term catheter users, the mean duration of use was 6 years (SD 7) and the median was 3.25 years. Therefore, patients and their families need to learn how to manage the catheter between nurse visits.

For short- or long-term catheter patients, the catheter should be the smallest size diameter that permits good urine flow. This should be 12-16 Fr for men and 12-14 Fr for women. Children’s sizes can range from 5-6 Fr on newborns to 5-10 Fr on toddlers to age 12. If a larger diameter was used to facilitate passage of blood clots, the size can be decreased by 2-4 Fr per catheter change until the optimal size is reached. Likewise, balloon sizes should be small, with 5- or 10-mL balloons, reserving the 30 mL only for bleeding episodes, usually postoperatively. Balloons of 2.5-5 mL can be used in children. Patients and their families should know the catheter and balloon size and the amount of water instilled to promote continuity of care. In a study of 202 persons, 8% did not know their catheter size, and 23% did not know the balloon size.

**Catheter Securement**

Patients who use long-term indwelling urinary catheters commonly use catheter securement devices to prevent urethral and bladder trauma and inadvertent catheter dislodgement. Post-hospitalization, securement is used to control postoperative bleeding or to protect a surgical anastomosis. Securing a catheter might also increase comfort and reduce CAUTI. Over time, traction of the catheter can cause erosion of the urethra, and in men this damage can spread to the length of the penis. Therefore, all long-term catheters should be secured—in women to the thigh and in men to the lower anterior abdomen or high on the thigh. Suprapubic catheters also should be secured. In a case report, a non-secured suprapubic catheter balloon made its way to the outside of the bladder and caused a urine-carrying fistula to the skin surface.

Although nurses might say they believe catheter securement is desirable, they may not always do it. In a survey of 82 nurses, including 8 continence specialists, 98% indicated they favor catheter securement, but in a previous prevalence study, only 4% had secured their patients’ catheters.

Catheter securement devices can be improvised or manufactured, including adhesive and non-adhesive devices. (See Table 1. Catheter Securement Devices.) Improvised devices are made with adhesive tape, sometimes with safety pins and rubber bands; however, tape can be irritating to the skin and hard to apply and remove. An early randomized clinical trial (RCT) with 59 patients showed that the manufactured device is easier to apply and remove and can stay in place 2 days longer than tape.

Adhesive-backed devices are available and these are particularly useful in short-term catheter use or for individuals prone to dislodgement. However, adhesive-backed devices also can irritate the skin and cause pain when removed. This should be an important consideration for geriatric patients or others with fragile skin. When using adhesive-backed securement devices, alcohol-based or other special wipes need to be used for removal to decrease skin damage.

Most non-adhesive devices use a stretch band and a Velcro locking system to hold the catheter to the thigh. While these devices have adjustability and can even be used with large/obese patients, they still need to be used according to the manufacturer’s instructions to be sure that blood flow is not restricted. Latex straps, which are used to hold the drainage bag in place on the leg, often twist and pull on the leg hair, and the latex can cause skin irritation or allergic reaction. Placing a thin cloth underneath the strap can minimize irritation. Nurses should instruct patients not to put any strap too tight and carefully monitor the skin condition, and this especially true for patients with peripheral vascular disease. Non-adhesive types also include holster-like devices hung from the waist or netting/cloth supporting bags, which can be ordered through urological supply companies online.

**Drainage Bags**

Urine drainage bags are available as leg bags, in sizes from 270-1000 mL, which permit more freedom in movement and travel. Large sized overnight bags (2000-4000 mL) are often used in the hospital or for people who are bedridden, e.g., at end of life. A “belly bag” has a 1000-mL capacity,
Table 1. Catheter securement devices

**Adhesive devices**

<table>
<thead>
<tr>
<th>Manufacturer</th>
<th>Product</th>
<th>Advantage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bard</td>
<td>StatLock® Foley Stabilization Device</td>
<td>Stable and easy to release catheter by opening the button</td>
</tr>
<tr>
<td>Dale Medical</td>
<td>Hold-n-Place® Adhesive Patch</td>
<td>Transparent and can be used up to 3 days; underlying skin is visible</td>
</tr>
<tr>
<td>Hollister</td>
<td>Horizontal Tube Attachment Device (HTAD)</td>
<td>Stable, and may be left in place up to 5 days</td>
</tr>
</tbody>
</table>

**Non-adhesive devices**

<table>
<thead>
<tr>
<th>Manufacturer</th>
<th>Product</th>
<th>Advantage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bard</td>
<td>Bard Catheter Leg Strap</td>
<td>Prevents skin irritation. Can be used for long-term catheterized patients. Are washable and can be reused. (Dale Medical has adjustable bands for very large patients)</td>
</tr>
<tr>
<td>Dale Medical</td>
<td>Hold-n-Place® Leg Band</td>
<td></td>
</tr>
<tr>
<td>Posey</td>
<td>Posey Catheter Tube Holder Straps</td>
<td></td>
</tr>
</tbody>
</table>

and clothing adjustments need to be considered. Since women do not stand to void, they might feel embarrassment in public toilet stalls as this bag is often emptied standing. This bag also is recommended only for people with normal bladder pressure (not those with flaccid bladders) as pressure is required for drainage. Privacy and covering the bags—even simply covering it with a bag or cloth—is important aesthetically and in promoting feelings of well-being. Hand dexterity should be considered as some drainage ports are easier to use, such as ones that twist or flip open. The bag should be kept off the floor and below the bladder level, but not more than about 12 inches to prevent excessive suction of the catheter eyes into the bladder mucosa. Connecting tubes can be used to attach the bag at the lower leg area and permit easier emptying, but regular attention is needed to prevent traction or twisting of the tubing in this position. Attaching a leg bag continuously
to the catheter and then adding a bag at night has not been studied, though some people advocate for this believing that the system is more “closed” this way.

Cleaning and reusing drainage bags, while not ordinarily recommended, is common practice in long-term catheter users in the U.S.; where-as changing it at least weekly is done in the U.K. In a recent U.S. study, over half of the 202 long-term catheter users, cleaned and reused drainage bags, using simple household solutions such as soap and water or vinegar and water. If using a dilute bleach solution (1 part household bleach to 10 parts water),24 care is needed to avoid splashing on the eyes, skin, or clothing.

Sometimes the catheter bag can become discolored, bluish purple (purple urine bag syndrome[PUBS]) because of bacterial colonization and conversion to tryptophan which produces indigo (blue) or indirubin (red) color in the urine. This can be embarrassing, and it can be eliminated by treating the offending organism6 or by changing the drainage bag when discoloration occurs.

**Catheter Insertion**

While there are no set guidelines for fixed intervals for catheter changes, other than for alleviating blocked urine flow or in relation to treatment for a UTI, most catheters are changed about once a month. Some people need them to be changed more often if encrustation occurs after 2-3 weeks or it becomes hard to remove if left too long. For people without problems, the interval can be gradually increased to 6-8 weeks.59 Careful observation of a pattern of “catheter life” to identify when it might block and then planning pre-emptively is advised.10

Inserting and removal of catheters in the home can be challenging because of working alone. For a trouble-free insertion, prepare all equipment ahead, including connecting the catheter to the drainage tubing. Use of sterile technique is required. Secure good lighting by moving lamps or asking a caregiver to hold a flashlight. Placing a female on her side can provide a better view of the urinary meatus, and it can minimize spasticity in the legs. For females after urine flows, insert 1-3 inches.14 Fill the balloon with sterile water, usually 7-8 mL in a 5-10-mL balloon. If the balloon is easily dislodged, such as with coughing, filling can be increased gradually to a maximum of 10-15 mL, depending on manufacturer guidelines. If the patient has a 50-mL balloon to control bleeding, do not fill it part way as it can fill asymmetrically and erode into the bladder.14 Decrease the size to a 5-10-mL balloon as soon as feasible.

For males, lubricate the catheter well by sliding the entire surface of the catheter through one hand with a lubricated sterile glove. Inflate the balloon when it has been inserted to the bifurcation, and then gently withdraw until resistance indicating that the balloon is in the bladder neck. If the catheter does not move in and out when gently moved, it can be “stuck” in the urethra, and it should be removed and replaced as it can cause severe trauma to the urethra within a short period of time which can result in CAUTI and/or scarring. In one retrospective case review of catheter trauma in 6 boys (aged 1 month to 16 years), balloon inflation while in the urethra caused significant damage,29 and lubrication and insertion to the bifurcation of the catheter was advised by the authors. If the catheter does not go in, spasticity at the entrance to the bladder can be the problem. Waiting, asking the patient to breathe deeply, and distraction (such as counting) can facilitate insertion. While removal of the catheter is usually not an issue, it can be, especially with suprapubic catheters which can develop a “deflation cuff” which interferes with easy removal, and is often painful. If this happens, the next time, after removing the water with a syringe, return about 1 mL water into the balloon. This will smooth out the cuff and make the removal easier.30 Sometimes a balloon will not deflate, and this requires medical assistance as guide-wires can be used to break up crystals in the balloon.

**Preventing Catheter Problems**

The key catheter problems are CAUTI, blockage, leakage, and dislodgement.5 31 There are no proven methods to prevent these problems, and long-term catheterized patients will undoubtedly have bacteria in the urine, often several flora which change over time.32

**Catheter Encrustations Causing Blockage or Leakage.**

Although irrigation is not recommended, they are often used in practice.5 51 In one study of 202 catheter users, 42% irrigated, many of whom (18%) did so once or more daily.4 Irrigations will not decrease CAUTIs, and the encrustations and sediment usually persist because of urea splitting bacteria—particularly *Proteus mirabilis*—which increase pH and causes calcium, magnesium and phosphorus minerals to precipitate into the urine.33 Unfortunately, a high pH (more alkaline) will persist as long as urea-splitting bacteria are present, therefore,
an often suggested method to lower urine pH (acidify) with vitamin C is not likely to be of benefit. However, increased fluid intake can dilute the urine and sometimes helps in decreasing blockage and premature catheter replacement even when the pH is high because the pH point of mineral precipitation is not the same as the urine pH.35 Bladder stones, often caused by the same minerals, need to be removed by cystoscopy, and if possible the offending organism should be eliminated through antibiotics. Nevertheless, bacteria living in biofilms and protected by a crystalline matrix are not easy to eradicate.36 Some people, with healthcare provider approval, might be helped by taking cranberry juice tablets because it keeps bacteria from adhering to the bladder lining,27 but research in catheterized patients is limited and evidence of cranberry juice efficacy in other populations is also unclear.38 Moreover, the juice may be unacceptable to patients over long periods of time.38

Leakage (bypassing of urine) can be minimized by proper positioning of the catheter so that traction is not pulling the balloon against the bladder opening, antispasmodic medication, and by treating constipation.39 Smaller size catheters and balloons also cause less irritation to the bladder sphincter and thus less leakage.

Paying Attention.

There is some research indicating that paying attention to fluid intake and preventing accidental dislodgment can be of benefit.40,41,42 Nurses can teach their patients to be more aware of their catheter position during transfers and instruct caregivers/family to keep the drainage bag from being run over by a wheelchair or pulled out when turning in bed. Noticing early signs of CAUTI could bring earlier treatment. Changes in the color or character of urine (odor, sediment) as well as malaise or weakness are the most common symptoms in long-term catheter patients.9 A pilot study teaching self-management, including optimal fluid intake and preventing dislodgement was effective, but further research is needed.41,42 Appropriate fluid intake levels vary, and while 30 mL/kg of body weight has been recommended,43 what is right for an individual should be mutually determined with the patient and provider.

In conclusion, home care nurses often manage care for patients with indwelling urinary catheters, many of whom have the device for years. Knowing catheter-related best practices is essential for their patients’ health.

References


31. Wilde MH, Brasch J, Getliffe K, et al. Study on the self-management of urine flow in long-term catheter users, quality of life in long-term urinary catheter users, and urinary drainage bag decontamination. Dr. Wilde is the author or coauthor of 37 articles, book chapters, abstracts, and other publications and has given many presentations on the subject of long-term urinary catheter care and management. She also sits on the review boards of several medical journals. She lives in Whitesboro, New York.


Mary Wilde, RN, PhD is Associate Professor in the School of Nursing, University of Rochester, Rochester, New York. Her current research interests focus on the self-management of urine flow in long-term catheter use, quality of life in long-term urinary catheter users, and urinary drainage bag decontamination. Dr. Wilde is the author or coauthor of 37 articles, book chapters, abstracts, and other publications and has given many presentations on the subject of long-term urinary catheter care and management. She also sits on the review boards of several medical journals. She lives in Whitesboro, New York.

Feng Zhang, RN, is enrolled in the MS/PhD program at the School of Nursing, University of Rochester in Rochester, prior to which he worked as an RN in Bramlage House of Meadowlark Hill Retirement Community (Transitional Services), Manhattan, Kansas. In May 2011, he received his BSc in Nursing at Washburn University, Topeka, Kansas. He has received several awards and honors, including the Loretta Lord Fellowship (2012-2013). Mr. Zhang lives in Rochester, New York.

Perspectives is an education program distributed free of charge to health professionals. Perspectives is published by Saxe Healthcare Communications and is funded through an educational grant from Dale Medical Products Inc. Perspectives’ objective is to provide health professionals with timely and relevant information on postoperative recovery strategies, focusing on the continuum of care from operating room to recovery room, ward, or home.

The opinions expressed in Perspectives are those of the authors and not necessarily of the editorial staff or Dale Medical Products Inc. The publisher and Dale Medical Products Inc. disclaim any responsibility or liability for such material. Clinicians are encouraged to consult additional sources prior to forming a clinical decision.
1. Catheters for short-term use:
   a. Include silver oxide coated types.
   b. Should not include use of coated latex catheters, such as those with PTFE coatings.
   c. Are used with a closed system.
   d. Are used for over 1 month.

2. Long-term catheter users:
   a. Have persistent urinary retention and no other alternatives
   b. Should irrigate the catheter to prevent CAUTI
   c. Need to use a large size overnight bag and avoid leg bags to prevent bacteriuria
   d. Are mostly people with severe incontinence

3. Pure silicone catheters:
   a. Have a smaller internal lumen and thus promote better urine flow
   b. Are softer than other catheter types
   c. Have been known to contribute to erosion of the urethra in men
   d. Can hold the water in the balloon longer than coated latex catheters

4. Catheter securement:
   a. Is not likely to be of benefit because patients don’t like it
   b. Contributes to excessive latex allergy
   c. Is expensive as the devices are single use and need to be changed daily
   d. Can prevent urethral or bladder damage

5. Drainage bags:
   a. Should never be covered
   b. Are available in a wide range of sizes with different types of emptying ports
   c. Evidence indicates that leg bags must be connected to overnight bags
   d. Tend to turn purple because of environmental conditions

6. When inserting a catheter in a male:
   a. Insert the catheter about 2 inches after urine is returned
   b. Removed if resistance is met at any time
   c. Inserted to the bifurcation
   d. Use a minimum of lubricant which could be irritating to the urethra

7. What percentage of tracheostomy patients has been reported as discharged to home with the tracheostomy in place?
   a. 12%
   b. 28%
   c. 41%
   d. 53%

8. What is a recommended tracheostomy tube cuff pressure?
   a. 10-15 mm Hg
   b. 16-18 cm H₂O
   c. 20-25 cm H₂O
   d. 28-32 mm Hg

9. Securement devices are commonly made of all of the following EXCEPT?
   a. Cloth adhesive tape
   b. Cotton twill tape
   c. Beaded chain
   d. Fabric with Velcro

10. Why are tracheostomy tubes in children more easily dislodged?
    a. The child moves a lot
    b. The tube is uncuffed and small
    c. Securement is difficult
    d. Ventilator tubing pulling on the tracheostomy tube

---

### Participant’s Evaluation Questions

<table>
<thead>
<tr>
<th>Question</th>
<th>Strongly Disagree</th>
<th>Strongly Agree</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Identify correct catheter and balloon sizes for adults and children.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2. Describe different types of catheter securement.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3. List the indications for tracheostomy in adults and children.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4. Identify the issues concerning care of the tracheostomy and tracheostomy tube for the patient at home.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**For immediate results, take this test online at**
[www.saxetesting.com](http://www.saxetesting.com)

*or mail to: Saxe Communications, PO Box 1282, Burlington, VT 05402 • Fax: (802) 872-7558 • www.saxetesting.com*

<table>
<thead>
<tr>
<th>Questions</th>
<th>Mark your answers with an X in the box next to the correct answer</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>A</td>
</tr>
<tr>
<td>2</td>
<td>A</td>
</tr>
<tr>
<td>3</td>
<td>A</td>
</tr>
<tr>
<td>4</td>
<td>A</td>
</tr>
<tr>
<td>5</td>
<td>A</td>
</tr>
</tbody>
</table>

Vol. 10, No. 1 /10