Use of wound dressings to enhance prevention of pressure ulcers caused by medical devices

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Abstract
Medical device related pressure ulcers (MDR PUs) are defined as pressure injuries associated with the use of devices applied for diagnostic or therapeutic purposes wherein the PU that develops has the same configuration as the device. Many institutions have reduced the incidence of traditional PUs (sacral, buttock and heel) and therefore the significance of MDR PU has become more apparent. The highest risk of MDR PU has been reported to be patients with impaired sensory perception, such as neuropathy, and an impaired ability for the patient to communicate discomfort, for example, oral intubation, language barriers, unconsciousness or non-verbal state. Patients in critical care units typify the high-risk patient and they often require more devices for monitoring and therapeutic purposes. An expert panel met to review the evidence on the prevention of MDR PUs and arrived at these conclusions: (i) consider applying dressings that demonstrate pressure redistribution and absorb moisture from body areas in contact with medical devices, tubing and fixators, (ii) in addition to dressings applied beneath medical devices, continue to lift and/or move the medical device to examine the skin beneath it and reposition for pressure relief and (iii) when simple repositioning does not relieve pressure, it is important not to create more pressure by placing dressings beneath tight devices.

Introduction
Pressure ulcers (PUs) traditionally occur on soft tissue and bony prominences that are exposed to pressure, shear and microclimate changes. However, PUs can also be caused by medical devices attached to or nearby the patient and in uncommon places such as the urethra, ears, upper back, neck and knees (1). Medical device related (MDR) PUs are defined as pressure injuries associated with the use of devices applied for diagnostic or therapeutic purposes wherein the PU that develops has the same configuration as the device. Pressure injuries result from a variety of reasons: the characteristic of the materials used to construct the device, difficulties in adjusting or securing it to the patient’s body, prolonged pressure in the same place, and pressure forces causing local oedema (2).

While the exact mechanical forces leading to the ulcerations from these devices have not been fully elucidated, it is logical

Key Messages
- MDR ulcers can occur from any medical device
- MDR ulcers can become full thickness pressure ulcers
- MDR ulcers are reasonably preventable with thin dressings under the device
- skin beneath a medical device must be inspected every shift
- faulty medical devices should be returned to the manufacturer
to believe that devices secured tightly to the patient’s body can be a source of pressure and if the patient develops oedema in the tissues after the device was initially secured, pressure can be increased. In addition, devices that increase sweating beneath them can be recognised as a source of skin microclimate changes.

Incidence of medical device related PU

While the development of PUs on the sacrum and heels has been tracked for decades, the presence of PUs beneath medical devices has only recently been identified. The incidence or facility-acquired rates of MDR PUs are not widely reported. However, many institutions have reduced their number of traditional PUs (sacral, buttocks and heel) and therefore the significance of MDR ulcers has become more apparent (Figure 1). The largest study found \((N = 104266)\), reported that medical devices led to 19.9% of ulcers on the ear, 14.3% of ulcers on the sacrum/coccyx, 10.2% of ulcers on the heel and 8.8% of ulcers on the buttocks (3). The specific devices involved were not included in the report of these findings. Reported incidences or agency-acquired rates of MDR PUs from the literature along with the known device leading to the ulceration are shown in Table 1.

Predictors of risk for MDR PU

The mere presence of a medical device (17,18) or the insertion site of the device (19) creates the risk for ulcers. However, there are certain devices and certain patients who are at increased risk. Ackland and colleagues (8) examined the records of 299 subjects in critical care who were wearing Philadelphia collars and were awaiting medical clearance of cervical spine injury. Cervical collar PU development was predicted by ICU admission \((P = 0.007)\), mechanical ventilation \((P = 0.005)\), the necessity for cervical MRI \((P < 0.001)\) and time to cervical spine clearance \((P < 0.001)\). Time to cervical spine clearance was the major indicator; such that the risk of ulceration increased by 66% for each additional day of wearing the cervical collar. Powers et al. (20) conducted a prospective, descriptive study of 484 patients wearing the Aspen collar. They also concluded that number of days with the cervical collar were found to be a significant predictor of skin breakdown \((P < 0.0001)\). The presence of braided or beaded hair can also contribute to PU development (20).

The ear has been commonly reported as a location for PUs (2,3,5,14,21). Some of the devices that encompass the ear are oxygen tubing and oxygen straps. Pulse oximetry devices are often clipped on to the ear helix or lobe in patients who are vasoconstricted and when finger probes are inaccurate. One study reported the actual pressure exerted on the ear lobe from ear pulse oximetry devices at 20.7 mmHg; the significance of this finding is unknown (22).

Incidence of PUs on the face and nasal bridge of patients using non-invasive positive pressure ventilation (NIPPV) masks has been reported as at least 17% (10,11). NIPPV masks can be difficult to fit to the patient’s face, and without proper fit, the device does not work correctly or not work at all. If the mask causes discomfort, the patient will often opt not to wear it (23). Treatment was curtailed in over 50% of patients using masks that caused discomfort (11). Oxygen delivery devices are the most common devices leading to pressure ulceration in children (24).

Patients at highest risk of MDR PU have been reported to be those with impaired sensory perception, such as neuropathy, and an impaired ability to communicate discomfort, for example, oral intubation, language barriers, unconsciousness or non-verbal state (5). Patients in critical care units typify the high-risk patient and they often require more devices for monitoring and therapeutic purposes. A secondary analysis of 2079 cases of patients in critical care, step down and surgical units concluded that patients with medical devices were 2-4 times more likely to develop a PU of any kind (2). Inspection of skin beneath medical devices may be cumbersome, because the medical device may be an essential component of the patient’s medical care. Pain from PUs beneath medical devices may also go unreported in unconscious patients. Another risk factor for PUs from medical devices is the tightness of securing method. Devices to support the airway (endotracheal tubes and tracheostomy tubes) (25) tend to be secured tightly. The presence of a medical device is the most frequently cited risk factor for PU development in children, with incidence rates as high as 50% (24). Devices associated with PU development are listed in Table 2.

Skin moisture due to diaphoresis or secretions under the medical device may macerate the skin, making it susceptible to PU formation (26). Moisture is probably a cofactor for their development, making the skin less resilient in the presence of secretions or other humidity. Oedema under the device stretches the skin, making it more fragile and prone to pressure injury. Even if the device initially fits properly, patients may develop oedema after securing of the device, thereby increasing the tissue tension (27). Blood vessels in oedematous deep tissue are compressed from the external pressure of the oedematous fluids, and oxygen transport from capillaries to cells is also impaired in oedematous tissue (2).

Medical devices can be made from a wide variety of materials, such as plastic, rubber, metals, plastics, composites, ceramics or silicone. Manufacturers’ improvements in design are often to improve comfort, but many devices are made of rigid materials or are secured with rigid material. The rigidity and inelasticity in the device or securing device causes friction, rubbing and increased pressure in tissues (10). Cervical collars are often very confining and lead to high occipital pressure (28). PUs may also arise because of the incorrect use, poor positioning or fixation of the equipment (29). Fixation or application of equipment using materials such as adhesive tapes or braces to adjust or secure the device may irritate susceptible skin, especially if oedema develops around the device (2). In addition, bad selection of poor equipment (17), incomplete evaluation of the skin (3) or problems with ill-fitting devices (5) may also be reasons to increase developing of these lesions.

Expert panel conclusions

In response to the increasing concern about MDR ulcers, the expert panel reviewed published data on devices and their prevention. The method used by this panel is described...
by Black et al. (30). The panel recognises that we are not an endorsed or elected body such as the National Pressure Ulcer Advisory Panel (NPUAP), the European Pressure Ulcer Advisory Panel (EPUAP) or the Pan Pacific Pressure Injury Alliance. The purpose of this article is to inform practitioners of our findings. Each of the statements that follow are written based on the methodology adopted in the recent PU guidelines (31), in that they have a strength of evidence and a supporting body of literature.

| Location | Example photograph | Prevention options |
|----------|--------------------|--------------------|
| Bridge of the nose from Non-invasive positive pressure device | Pretreat the bridge of the nose with dressings prior to application of mask |
| Nasal ala from NG tube | Splint NG tube to suspend it from ala |
| Elastic stocking rolled down by patient because it was too tight | Use proper size of stockings, remove twice daily for an hour and inspect the skin beneath stockings |
| Lip from ET tube | Rotate ET tube on every shift or more often (such as every 2 hours with turning) |
| Occiput from neck collar | Remove hard collars as soon as possible and replace with softer collar. Release braided or beaded hair prior to fitting with collar if possible. Examine scalp daily |

Figure 1 Examples of medical device-related pressure ulcers.

Consider applying dressings that demonstrate pressure redistribution and absorb moisture from body areas in contact with medical devices, tubing and fixators [strength of evidence (of recommendation), (SOE) = B]. Three studies were found that reported on the effect of dressings placed between the device and the patient’s skin. Weng (12) conducted a quasi-experimental study with three groups of patients using NIPPV masks comparing time-to-nasal ulceration with two dressings and a control group without
Table 1 Type of device and frequency of ulceration

| Device                          | Frequency (Reference)                                                                 |
|---------------------------------|---------------------------------------------------------------------------------------|
| Cervical neck collar            | 7.2% (4); 22% (5); 23.7% (6); 33% in 5 days, 44% over 5 days (7) Ulceration from neck collar increased 66% for every 1 day in device (8) |
| Endotracheal tube               | 10.5% (6)                                                                             |
| Immobilisers/Halo casts         | 17% (5); 4% (9)                                                                       |
| Nasogastric tubes               | 8% (5)                                                                                |
| Non-invasive positive pressure  | 17% (10,11)                                                                           |
| ventilation devices             | 96.7% (12)                                                                            |
| Oxygen tubing                   | 12.9% in children (13); 13% (5); 19.9% (3)                                            |
|                                 | 28–47% (14)                                                                           |
| Pulse oximetry monitors         | 9% in children (15)                                                                   |
| Stockings or boots              | 12% (5)                                                                               |
| Tracheostomy flange and ties    | 8.1% (16)                                                                             |

dressings. One group had a film dressing, another group had a hydrocolloid dressing, and another was a control with no dressing. Ulceration rate was 96.7% in the control group, 53.3% in the group with film dressings and 40% in the group with hydrocolloid dressings. The skin was inspected every 30 minutes and the time-to-ulceration differed between groups with the shortest time in the control group (1111 minutes) and longest time in the hydrocolloid group (3272 minutes). Boesch and colleagues (16) reported a decrease in PU incidence from a baseline of 8.1% to 3.4% after the introduction of a bundled change programme including education and Mepilex® Lite™ (Mohnlycke Health Care; Gothenburg, Sweden) dressing under the tracheostomy tube flanges. Turgania et al. (14) reported a reduction in the incidence of ulcers on the ear due to nasal cannula from 37% to 0% with the use of padding on the ears. A comparison of foam dressings to other types of dressings has not been reported; however, the attributes of foam dressings would appear to have benefit in these patients by reducing pressure and absorbing moisture, where hydrocolloids and films cannot absorb moisture.

Other studies examined products to hold the medical device away from the body orifice. Bite blocks were used to prevent oral PUs (32), splints for nasotracheal tubes (33) and silicone paste moulded to the bridge of the nose (11). The incidence of PUs beneath tracheostomy tubes or ties and/or the connection to the ventilator dropped to zero after the tracheostomy tube length was extended (16).

In addition to applying dressings beneath medical devices, continue to lift and/or move the medical device to examine the skin beneath it and reposition for pressure relief (SOE = C)

This recommendation is from expert opinion. It is appreciated that the use of a dressing on the skin beneath a medical device is not a replacement for other prevention strategies. Skin should be inspected according to agency policy or accepted standards of care. Skin must be inspected for signs of pressure due to the device, which cannot be seen while the device is still

Table 2 Medical devices associated with pressure ulcer formation

| Medical device              | Anatomical site                   |
|-----------------------------|-----------------------------------|
| Endotracheal tubes          | Lips                              |
|                            | Tongue                            |
| Nasotracheal tubes          | Nose                              |
|                            | Nares                             |
|                            | Nasal bridge                       |
| Nasal cannulas             | Ears                              |
| Oxygen tubing              | Nose                              |
| Cervical collars           | Neck                              |
|                            | Occiput                           |
|                            | Clavicle                          |
|                            | Chin                              |
|                            | Mandible                          |
| Oximetry probes            | Fingers                           |
|                            | Ears                              |
|                            | Nose                              |
| NIPPV/BIPAP                | Forehead                          |
|                            | Nose                              |
| Tracheostomy tubes         | Front of the neck                 |
| Tracheostomy braces        | Around/back of the neck           |
| Stockings/compression devices | Behind knees                    |
|                            | Achilles tendon                   |
|                            | Lower extremities                 |
| Immobilisers               | Ankles                            |
|                            | Wrists                            |
| Faecal containment         | Perianal                          |
|                            | Buttocks                          |
| Foley catheters            | Urethra                           |
|                            | Upper thighs                      |
|                            | Buttocks                          |
|                            | Perianal                          |
|                            | Heels                             |
|                            | Arms                              |
|                            | Elbow                             |
|                            | Ankles                            |
| Abdominal binders          | Wristis                           |
|                            | Neck                              |
|                            | Ribs                              |
|                            | Skin folds                        |
| External fixators          | All locations where the device is in contact with skin |
| Halo rings                 |                                   |
| Vest brace                 |                                   |
| Catheters                  |                                   |
| Tubing                     |                                   |
| Orthotics                  |                                   |
| Drains                     |                                   |
| Stopcocks                  |                                   |
| G/J tubes                  |                                   |
| JPs                        |                                   |
| T-tubes                    |                                   |
| Ostomy clips               |                                   |
| Bedpans                    |                                   |
| BIPAP, bi-level positive airway pressure; NIPPV, non-invasive positive pressure ventilation. |
in place. For example, an endotracheal tube must be moved to inspect the lips for pressure injury. In obese patients, the medical device may become hidden it the skin folds, and the nurses need to be diligent to account for all medical devices in these patients. Three-quarters (74%) of device-related PUs were not identified until they were stage III, stage IV or unstageable compared with 54% of non-device-related PUs. Twenty percent of non-device-related PUs were first identified when they were stage I, compared with only 5% of device-related ulcers (5). The dressing must be removed to visualise the skin, and then reapplied before replacing the medical device. An important consideration is the ability of dressing to be removed without injury to the skin during repeated assessments. The bridge of the nose is thinly covered with skin and could be easily injured with highly adhesive dressings. If possible, avoid placing a medical device back onto pressure-injured skin.

More than 50% of the medical devices that contributed to these pressure-related skin injuries were SpO2 probes, artificial airways or bi-level positive airway pressure (BiPAP) masks. The mechanism of device-related pressure injury is similar to that of PUs caused by immobility, but prevention differs. Prevention includes rotating SpO2 probe sites, avoiding securing the endotracheal tube close to the oral commissure, and inserting pressure-relieving material between oxygen masks and the nasal bridge (34).

Agencies may have to determine which member of the team, for example nurses or respiratory therapists (or both) are best suited to move devices and inspect the skin beneath them. Some medical devices are not meant to be moved, such as a cast. In those situations, follow accepted practice for care of patients when skin cannot be seen beneath the device. Agencies will also have to assist the staff to determine the proper size for a device by placing product selection guides in patient-care areas. Support stockings are often applied prior to surgery, and then become too tight when the limb becomes swollen after surgery. Similarly, some medical devices are applied prior to fluid resuscitation (e.g., sepsis) and the securement product, such as tape or straps, becomes tighter as fluid is shifted into interstitial spaces. The early signs of pressure from the medical device and the securement device should be identified and adjustments made in the device or straps to prevent injury.

When simple repositioning does not relieve pressure, it is important not to create more pressure by placing dressings beneath tight devices (SOE = B)

Two studies were found that reported on the effect of thin dressings placed between the device and the patient’s skin. Weng’s (12) quasi experimental study reported that the use of even a thin (film/hydrocolloid) dressing applied to the nose reduced the incidence of nasal bridge PUs by 43-4% and 56-7%, respectively compared to an ulceration rate of 96-7% when no dressing was used. Boesch and colleagues (16) also saw a 4-7% decrease in ulcers under tracheostomy flanges with a bundled change programme including Mepilex® Lite™ under the tracheostomy tube flanges. Gunlemez et al. (35) reported that the use of silicone gel sheeting dressings reduced MDR ulcers due to oxygen delivery devices (P < 0-05).

In these studies, thin dressings appear to reduce the magnitude of the pressure. Inserting thick dressings for prevention has not been reported, but the authors offer the opinion that more pressure could be created with a thick dressing.

Reporting MDR ulcers

Root cause analyses in these cases indicated that staff were sometimes uncertain about whom to contact about an ill-fitting device, were unsure of what a well-fitted device should look like against the skin, or were not aware of the potential for oedema to affect fit of devices or risk of PU development (2,5). Common characteristics included impaired sensory perception, such as neuropathy, and an impaired ability for the patient to communicate discomfort, for example, oral intubation, language barriers, unconsciousness or non-verbal state. The lack of clarity around best practices for skin inspection with the use of devices may have been a contributing factor in device-related PUs not being identified until they had become more advanced (5,36). Wounds from medical devices are PUs; they should not be classified as wounds from underlying disease states such as arterial insufficiency or be overlooked in institutional reporting. MDR ulcers should be reported separately from other PUs (those on the sacrum, buttocks and heels, for example and not associated with the use of a medical device in the same body area). Further policy work is needed on how to report PUs from medical devices on the mucous membranes that leave no scar and heal quickly. These wounds are PUs but do not appear to cause permanent damage to the patient. A stratification system on severity or likelihood of healing for MDR ulcers would aid in the analysis and reporting issues. Facilities required to do root or common cause analyses on every ulcer can find the process burdensome for some of the quickly healed ulcers on mucous membrane. It is important not to mistake the idea of a lack of scar to equate to an insignificant injury. Any open wound can worsen and methods to reduce the number of these wounds should not be overlooked.

The name of the device that led to the PU should also be recorded in order to keep track of the device causing harm. For example, it may not become apparent that the same medical product is causing harm until the data from the entire hospital system are reviewed. The authors believe that when a pattern of MDR PUs is apparent, the information should be conveyed to the manufacturer and regulatory bodies. If hospital staff continually accepts the development of MDR ulcers as a quality improvement problem, and never report them to the manufacturer, opportunities for improved safety profiles in product design are missed.

Conclusion

PUs from medical devices are increasing in frequency, and it has been said MDR PUs are an epidemic. High-risk patients should be identified during risk assessments, skin inspection beneath devices must occur at every shift and the devices
should be moved as much as possible (for example moving the endotracheal tube and pulse oximetry probe from side to side with each body turn). Applying protective dressings to pad the skin is imperative; most of these ulcers are completely preventable. However, medical professionals can only work with the devices provided. The solution also depends on the development of safer materials that do not increase the risk of developing a PU.

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