Recognising, Managing and Preventing Deep Tissue Injury (DTI)

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Recognising, managing and preventing deep tissue injury (DTI)
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This document was developed by the Expert Working Group with the objectives of:
- Raising awareness and understanding of the diagnosis, management and prevention of deep tissue injuries (DTIs)
- Encouraging communication across all healthcare settings, including the community, where there are patients at increased risk of DTI.

In doing so, however, the Group recognises that much remains to be learnt about DTI. Development of the document began with a day-long meeting and was followed by extensive review of the resulting text before production of this final document.

By explaining current understanding of the mechanisms involved in the development of DTI and describing how to recognise, treat and prevent DTI, the document aims to improve diagnosis, management, prevention and reporting of DTIs in the UK.

The chart ‘Deep tissue injury (DTI) - an overview’ (page 1) summarises the content and guidance provided in this document and links to the appropriate pages.

**Key**

- Key practice point
- Key recommendation and call to action
Deep tissue injury (DTI) - an overview

What is a DTI?
Pages 3-5

A DTI is a type of pressure ulcer (PU); occurs most commonly at the heels, sacrum and buttocks (including the ischial area)

- In light-toned skin often presents as an area of non-blanching purple or maroon skin and in dark-toned skin, as discolouration or lighter/darker skin
- May feel firmer or softer (‘boggy’ or ‘mushy’) and warmer or cooler than surrounding tissues
- 24–48 hours later, the area may look dry; the epidermis may peel or a shallow blister may form that may or may not break open
- Blisters can be filled with a dark liquid and look like blood blisters; edges of the blister will be non-blanching
- May develop a hard, blackened surface (eschar); when this sloughs or is debrided a deep wound may be revealed
- May deteriorate rapidly to an open wound, resolve before the skin breaks or remain static

What causes a DTI?
Pages 6-8

A DTI results from external application of a mechanical load (pressure), to the skin and subcutaneous tissues over a bony prominence such as the heel

- The load causes tissue necrosis by producing direct structural damage to cells and by reducing blood flow and drainage of fluids and waste products. Return of blood flow may add to the tissue damage (reperfusion injury)
- DTIs start at the bone: tissue interface and develop from the ‘inside out’
- As a result, a DTI may not become apparent for 24–72 hours after the mechanical loading event that caused it

Who is at risk of a DTI?
Pages 9-10

DTIs share many risk factors with PUs, e.g. restricted mobility, immobility, altered posture, poor blood flow to tissues due to heart or lung disease, smoking, diabetes, reduced sensation, incontinence, increased age, poor nutrition and poor general health

- Factors particularly increasing risk of a DTI are restricted mobility, immobility, impaired sensation and reduced tissue tolerance
- Patients at increased risk include those with a sudden unexpected period of reduced mobility due to collapse, fracture or cerebrovascular accident, those on an emergency department trolley or undergoing prolonged surgery, those receiving intensive care, acute care or long-term care, and those that are transferring between facilities or transitioning from being bed bound to sitting in a chair

How is a DTI assessed?
Pages 11-20

Several conditions other than DTI present with discoloured, purple or maroon skin

- Exclude Category 1 or 2 PU, bruise, haematoma, blood blister, skin tears and incontinence-associated dermatitis before diagnosing a DTI
- Conduct assessment, management and prevention of DTI in the context of the principles of mental capacity and consent
- Even if the diagnosis is unclear, refer as necessary and at the same time treat as for a DTI
- Undertake formal assessment, including full skin inspection and PU risk assessment
- PU risk assessment tools should be used alongside skin inspection and clinical judgement
- Use a structured wound assessment to assess the DTI
- Document assessment; report according to local guidelines for incident reporting

How is a DTI managed?
Pages 21-26

Define objectives, remembering that sometimes other care needs may take priority

- Optimise patient condition: nutrition and hydration, management of comorbidities
- Offload the DTI, i.e. ensure no contact with a support surface and no mechanical load
- Reposition regularly and use pressure redistribution support surfaces
- Manage pain, including pain due to repositioning or wound care
- Keep patient clean and dry; moisturise dry skin
- If the skin is broken or there is an open wound, cleanse and cover with an appropriate dressing
- Monitor and regularly reassess
- Refer as appropriate, e.g. to a tissue viability service, if the DTI deteriorates rapidly or has not improved within two weeks
- Educate the patient, family and carers
- Document objectives and management strategies

How is a DTI prevented?
Pages 27-28

Identify patients at risk by recognising risk factors (see above ‘Who is at risk of a DTI?’), using formal PU risk assessment tools and regularly inspecting skin

- Clinical judgement should be used in combination with PU risk assessment tools
- When a patient is recognised to be at risk of DTI:
  - Perform a holistic assessment
  - Reduce risk of tissue damage, e.g. offload, implement regular repositioning and use pressure redistribution support surfaces, while considering all of the patient’s needs
  - Optimise patient condition: nutrition, management of comorbidities, pain management
  - Educate the patient, family and carers
  - Monitor and reassess risk of pressure damage and effectiveness of prevention regimen
- Document assessment and prevention strategies
The challenges of DTI

A deep tissue injury (DTI) is a type of subcutaneous tissue damage that results from an externally applied mechanical load (pressure). Recognition that DTIs have features distinct from pressure ulcers (PUs) led to DTIs being added in 2009 to the international classification of pressure ulcers (PUs) produced by the European Pressure Ulcer Advisory Panel and National Pressure Ulcer Advisory Panel (EPUAP/NPUAP, 2009). A DTI is not always associated with broken skin and does not always become the open wound often defined as a PU. However, DTIs do have the potential to develop into large, deep wounds with significant tissue loss that have serious consequences for patients and healthcare systems (Peart, 2016).

The formalisation of DTIs as a distinct category within PU classification is relatively recent. This, along with recent changes in PU terminology in some parts of the world (Box 1), and variations across the UK’s healthcare systems in requirements for reporting and data collection, has caused confusion. It is likely that DTIs are under-reported because they are not always easy to recognise, particularly in the early stages, and may be miscategorised or not reported. In addition, some reporting systems do not include DTI or delay recording of the tissue damage until classification as one of the main PU categories is made.

Box 1. Terminology issues

Suspected deep tissue injury (sDTI) or deep tissue injury (DTI)?
When DTIs were first included in the international classification for PUs, the term DTI was preceded by the word suspected (abbreviated to sDTI) partly because of a lack of understanding of the condition and its relationship to other types of pressure-related damage (EPUAP/NPUAP, 2009). However, the pathophysiology of DTIs is now much better understood, and it is recognised that externally applied mechanical loads, such as pressure and shear, can give rise to DTIs and PUs in different ways (Oomens et al, 2015). For this reason, and because of other concerns over the use of the word, the recently updated PU classification produced by the NPUAP has dropped the ‘suspected’ preceding DTI (Edsberg et al, 2016).

The Expert Working Group also proposes that ‘suspected’ is omitted before DTI in the UK to remove any misunderstanding about the existence of DTI as a distinct condition and to encourage appropriate reporting, management and prevention.

Pressure ulcer (PU) or pressure injury (PI)?
In 2016, the NPUAP decided to replace the term ‘pressure ulcer’ (PU) with ‘pressure injury’ (PI) (Edsberg et al, 2016). This brought NPUAP terminology in line with that used in countries covered by the Pan Pacific Pressure Injury Alliance (PPPIA) (Australia, Singapore, Hong Kong and New Zealand) (AWMA, 2012). Reasons for this change included the desire to emphasise both the preventable nature of many PUs and the fact that not all PUs represent open wounds. However, there are concerns with the use of the term ‘injury’ because it implies a very acute event leading to the tissue damage, even though it is known that the damage associated with PUs may take some time to develop (Bader & Schoonhoven, 2016; Ousey et al, 2017a). Indeed, the time element underpins the rationale for repositioning in the prevention of PUs. In addition, there are concerns that using the word ‘injury’ might be seen to imply that healthcare providers have caused the injury; a concern that has also been raised in the US (Mrdjenovich et al, 2016; NPUAP Position Statement, 2017). Finally, replacing the word ‘ulcer’ removes alignment with other wounds such as diabetic foot ulcers (DFUs) and venous leg ulcers (VLUs). This is particularly unfortunate in the case of DFUs, which often share some of the same aetiological mechanisms as PUs (Vowden & Vowden, 2015).

In common with a recent NHS Improvement consensus meeting that voted to use ‘pressure ulcer’ in England, the Expert Working Group recommends retaining the term ‘pressure ulcer’ (PU) in the UK.
Defining DTI

DTI is a type of subcutaneous tissue damage, affecting muscle where present, that originates close to bone and is the result of a mechanical load, such as pressure, applied to the skin. Box 2 contains the definition of DTI from the most recent (2014) NPUAP/EPUAP/PPPIA PU prevention and management guidelines. As noted in Box 1 (page 2), the Expert Working Group recommend using ‘deep tissue injury’ and omitting ‘suspected’.

RECOGNISING DTI

The appearance of a DTI on presentation depends on when it is recognised in the time course of DTI development. As the name suggests, DTI starts deep within tissue and does not usually become apparent until about 24–72 hours after the event that caused the tissue damage (Black et al, 2016). Such events include lying immobile on a hard surface, e.g. on the floor after a fall that has resulted in unconsciousness or that is due to a stroke, or experiencing a period of poor tissue perfusion, e.g. during cardiothoracic surgery. In keeping with pain being a predictor of PUs (Smith et al, 2017), conscious and sensate patients may complain of pain before the appearance of the physical signs of DTI. See pages 9-10 for more detail on risk factors for DTI.

In patients with light-toned skin, DTI presents about 24–72 hours after initiation as a demarcated area of purple or maroon skin, often likened to a bruise. The purple or maroon area may be surrounded by an area of erythema (Figure 1). The discoloured area and erythema are non-blanchable (Box 3, page 4) (Black et al, 2016).

When palpated, the affected area may feel:
- Firmer or hardened (indurated) due to stiffening of dead or dying muscle tissue or
- Softened (‘boggy’ or ‘mushy’) as the damaged tissue starts to break down and/or
- Warmer or cooler than surrounding areas (Gefen, 2009).

In patients with dark-toned skin, the skin of affected areas may be darker or lighter, and palpation to detect changes in tissue texture or temperature is particularly important (Black et al, 2016). A retrospective review of patients with darkly pigmented skin found that the most commonly described presentation of DTIs was purple discolouration of intact skin (Sullivan, 2014).

Patients who are conscious and have sensation in the affected area may complain of discomfort or pain, and/or find palpation uncomfortable or painful

When assessing the temperature of the skin, it is important to consider whether the area has been in contact with or covered by bedding or clothing or has been exposed (i.e. it is warmer or cooler for extrinsic reasons), the ambient temperature, and the temperature of the assessor’s hands.
It is important to palpate the skin and subcutaneous tissues of any area that may have tissue damage.

After a further 24–48 hours, the outer layer of the skin of the affected area may appear dry, start to peel or form a shallow blister (known as epidermal lift or sloughing). The full thickness of the skin may also lift or blister, and breaks in the skin can occur to reveal a purple, maroon, black or white wound bed (Black et al, 2016). In patients with dark-toned skin, blistering and skin breaks may be the first signs noticed of a DTI (Sullivan, 2014). Blisters may become filled with liquefied necrotic tissue and look like blood blisters.

Eventually a hard, blackened surface (eschar) may develop. When this finally sloughs or is debrided it will reveal the extent of the tissue damage and possibly a deep open wound. In some cases, sloughing of the eschar reveals intact or healed skin.

Although the appearance of DTI is often described as a bruise, it should be noted that a DTI does not go through the colour changes typical of a bruise as it resolves.

NATURAL HISTORY AND EVOLUTION OF DTI

The potential evolution of a DTI is illustrated in Figure 2, page 5. Although full clarification of the natural history of DTIs and the factors that influence it is still to be established, it is clear that the outcome is variable. Some DTIs can deteriorate very rapidly to become large, deep open wounds despite best practice management. However, not all DTIs evolve or deteriorate. A proportion of DTIs will remain static for quite some time (anecdotally up to several months) or will resolve without forming an open wound (Richbourg et al, 2011).

In the experience of the Expert Working Group, when resolution occurs it is often within 7–10 days of diagnosis of the DTI.

Currently, it is not known what proportion of DTIs resolve or whether there are indicators that could be used to predict which DTIs will resolve. A two-year study that followed 128 DTIs for an average of 6 days found that 37.5% of DTIs had resolved at the end of follow-up (Sullivan, 2013).

DTIs that resolve or remain static need continued care and monitoring to ensure the area is not exposed to further mechanical loading and to prevent repeated damage or extension of the injury.

Box 3. Testing for non-blanchable erythema (NPUAP/EPUAP/PPPIA, 2014)

Gentle pressure is applied to the discoloured or reddened skin for three seconds using either a finger or a clear plastic device.

a) Blancheable erythema – application of pressure will push blood away from the area that has been pressed turning it paler than the surrounding skin (see photograph).

b) Non-blanchable erythema – if the blood flow to the area being tested has been damaged, the application of pressure will not make the skin become paler and it will remain discoloured or reddened. Non-blanchable erythema indicates tissue damage.
LOCATION OF DTI

An analysis of US data from a large international pressure ulcer prevalence survey undertaken in 2006–2009 found that DTIs accounted for about 9% of all PUs. The study also found that 41% of the DTIs occurred at the heel, 19% over the sacrum and 13% on the buttocks (VanGilder et al, 2010) (Figure 3).

A more recent analysis of hospital-acquired DTIs recorded over a two-year period (2010–2012) in one US hospital, reported the coccyx, heels and buttocks to be the most commonly affected sites. This study also reported DTIs to have occurred in relation to medical devices, and at the sacrum, intergluteal area, trochanter and ischium (Tescher et al, 2017).

The heels are thought to be particularly at risk of DTI because the underlying bony prominence (the calcaneus) has a small radius of curvature, i.e. is relatively pointed, and the layer of tissue overlying the bone is relatively thin (Salcido et al, 2011). This means that the intensity of a mechanical load applied to the heel can be high.

The location of a DTI will depend on the position of the patient. For example, in a patient who is sitting upright in a chair, DTI may occur under the ischial tuberosities, and in a patient who is sitting in bed with the head of bed elevated, DTI may occur over the sacrum. Similarly, a patient who has been lying on the floor after a fall or collapse due to acute illness may have evidence of damage at points that have been in contact with the floor. DTI may also occur under medical devices such as casts, splints, oxygen face masks or elastic bandaging (Tescher et al, 2017).

DTIs are most commonly found at the heels, sacrum and buttocks, but may occur in any location where prolonged pressure is applied, including under medical devices.

As DTIs can occur at anatomical sites where PUs may occur, pressure-related damage, including any related to a medical device, should be carefully assessed to determine whether the damage is a DTI or a PU.

Figure 3: Sites of DTI occurrence (VanGilder et al, 2010)
PUs and DTIs occur because of tissue damage caused by the application of a mechanical load (Box 4) to the skin, subcutaneous tissues and muscle. In general, higher mechanical loads will result in tissue damage more quickly than lower loads (Figure 4).

**Box 4. Mechanical load, pressure and shear in the context of PUs and DTIs (International Review, 2010; Gefen et al, 2013)**

- **Mechanical load** - a generic term that covers all forces, including pressure and shear, applied to the skin and subcutaneous tissues, e.g. muscle
- **Pressure** - results from the application of a force perpendicular (i.e. at right angles) to the surface of the skin. The pressure compresses the tissues and can distort or deform the skin, subcutaneous tissues and muscle. Tissue distortion is likely to be greatest when pressure is applied over a bony prominence
- **Shear** - causes layers of body tissues to move relative to each other and may occur:
  a) Superficially - e.g. as a result of a force applied parallel (tangentially) to the surface of the skin
  b) More deeply (internally) - e.g. as the result of deformation of skin and muscle when pressure is applied over a bony prominence

**Figure 4: Effects of magnitude of mechanical load and time on likelihood of tissue damage (adapted from Stekelenberg et al, 2008; Gefen et al, 2008; International Review, 2010)**

**Threshold for tissue damage:**
- ✔ higher risk for tissue damage occurs in the area above the line
- ✗ lower risk for tissue damage occurs in the area below the line
- ⬤ Higher mechanical loads reach the threshold for tissue damage earlier than lower mechanical loads

**MAIN MECHANISMS**

A mechanical load can have several effects that may ultimately damage tissue and cause tissue breakdown and cell necrosis (Figure 5, page 7). The two main mechanisms involved are:

- **Structural damage** - The mechanical load has a direct effect on tissue cells, damaging cell membranes and disrupting internal cell structure to cause cell death. In vitro and animal studies have shown that this type of damage can occur within tens of minutes of application of the mechanical load and certainly much more rapidly than the damage caused by ischaemia (Gawlitta et al, 2007; Oomens et al, 2015).
- **Ischaemia** - The mechanical load can compress capillaries causing partial or complete occlusion. This reduces the delivery of oxygen and nutrients to tissues. If the rate of delivery is below the physiological demands of the tissues, ischaemia will occur. If the ischaemia is sustained for long enough and/or at high enough levels, tissue death can occur due to a change in metabolism and the accumulation of waste products. Generally, ischaemia takes a few hours to produce tissue damage, and muscle is more susceptible to ischaemia than skin (Agam & Gefen, 2007; Oomens et al, 2015).
The amount of tissue deformation, i.e. the degree of change in tissue shape in response to the mechanical load, required to produce ischaemia is lower than that required for structural damage (Oomens et al, 2015). However, the threshold at which damage occurs from either mechanism will vary from patient to patient and will depend on numerous factors, e.g. the presence of comorbidities (such as cardiovascular disease and diabetes) that may reduce tissue perfusion and the condition and thickness of the tissue layers between the skin and bony prominence (Oomens et al, 2015). A combination of reduced tissue perfusion and less tissue between the skin and bony prominence may result in DTI particularly quickly.

OTHER EFFECTS INVOLVED

Two other mechanisms to consider in the aetiology of DTIs and PUs are:

- **Lymphatic occlusion** – In addition to occluding blood vessels, mechanical loads can occlude lymphatic vessels and may cause the accumulation of metabolic waste products that contribute to tissue damage (Miller & Seale, 1981; Peart, 2016).

- **Ischaemia-reperfusion injury** – Although counterintuitive, restoration of blood flow to an ischaemic tissue can exacerbate ischaemia-induced damage. Reperfusion initiates an inflammatory response that stimulates the formation of oxygen-derived free radicals and other agents that can cause further tissue damage (Collard & Gelman, 2001). An animal study of DTI found that reperfusion reversed tissue damage after a limited time of ischaemia, but that if ischaemia was prolonged, reperfusion exacerbated the existing tissue damage (Loerakker et al, 2011). Patients who experience hypotension, whether as a consequence of disease or as a result of other treatments, may be particularly prone to the effects of ischaemia-reperfusion injury.
DIFFERENCES IN THE DEVELOPMENT OF DTIs AND PUs

The differences in clinical progression of DTIs and PUs suggest that although the underlying causes are similar, the initial site of damage is different. In DTIs, mechanical loading in the form of externally applied pressure causes tissue damage that occurs initially adjacent to the bony prominence and in the muscle layer if present (Figure 6). The damage then moves towards the surface in a manner sometimes described as ‘bottom up’ or ‘inside out’.

Conversely, in PUs, the mechanical load comprises pressure with superficial shear. These loads cause tissue damage initially at the surface; the damage works downwards, or ‘top down’ or ‘outside in’ (Fife, date unknown; WUWHS, 2016).

In another analogy, DTIs have been likened to geological sinkholes. This analogy is also useful because it is in keeping with the frequent finding in DTIs of undermining due to tissue damage that is more extensive than initially apparent from the skin surface. In a similar analogy, PUs have been likened to potholes in a road (WUWHS, 2016).

Although a clear distinction has been made here to aid understanding of the differences in the development of DTIs and PUs, in clinical practice it is possible that the mechanisms sometimes occur together in the development of an area of pressure-induced damage. For example, a patient who has been immobile may start to develop a DTI, but then may regain some movement that introduces superficial shear which also contributes to the tissue damage.

The facts that DTIs can occur very quickly and start below the skin surface (often in muscle) suggest that high mechanical loads may be involved and that structural damage to cells is a major cause of tissue necrosis. These facts reinforce the need for timely risk assessment and frequent skin inspection.
An individual’s risk of developing a DTI or a PU is dependent on a complicated interplay of numerous factors (Coleman et al, 2013). At a basic level, the risk is related to the intrinsic ability of the patient’s tissues to withstand the effects of extrinsic risk factors such as externally applied mechanical loads, i.e. to a level of tissue tolerance. When the effects of the extrinsic risk factors exceed the specific tissue tolerance, the individual is at an increased risk of developing a DTI or PU (WUWHS, 2016). Tolerance to extrinsic risk factors will be reduced by a range of intrinsic risk factors (Box 5, page 10) such as immobility, reduced tissue perfusion and reduced sensation.

FACTORS THAT INCREASE THE RISK OF DTI
PUs and DTIs share many risk factors, but there are some risk factors that particularly increase the risk of a DTI. These can be divided into factors that restrict immobility or cause immobility, impair sensation, and reduce tissue tolerance (Figure 7, below; and Table 1, page 10).

In some conditions, several of these factors may be present and contribute to increased risk of DTI. For example, individuals with spinal cord injury can have reduced mobility, impaired sensation, hypotension and muscle atrophy (wasting) and so are at an increased risk of DTI (Gefen, 2014). A recent analysis of factors contributing to DTI in intensive care unit patients reported that the odds of developing a DTI increased by 20% for every hour increase in length of surgery (Kirkland-Kyhn et al, 2017).

Patients in intensive care units (ICUs), undergoing surgery, receiving acute care in hospital or in long-term care are at increased risk of DTI (Fleck, 2007)

Patients undergoing transitions in care, e.g. when being transferred from one facility to another (Preston et al, 2017), can be at an increased risk of DTI, particularly if not placed on an appropriate support surface during transfer or in the receiving clinical area

Patients undergoing other transitions, e.g. from being bed-bound to sitting in a chair, are also at an increased risk of DTI because the change in position will change the ways and the time period that mechanical loads are applied to the tissues and because the support surface may not be appropriate

Patients that have had a DTI may be at increased risk of the development of further DTIs or PUs because during the healing process the tissues are likely to have been remodelled, e.g. with scar tissue, and to have altered physical characteristics.

Who is at risk of DTI?

![Figure 7: Categories of factors that increase risk of DTI](image-url)
Box 5. Main intrinsic risk factors for PU and DTI (AWMA, 2012; Coleman et al, 2013; Coleman et al, 2014)

**Major intrinsic risk factors**
- Immobility/reduced activity/altered posture – e.g. bed-fast, chair-fast, contractures
- Reduced tissue perfusion – e.g. due to peripheral vascular disease, cardiopulmonary disease, diabetes, smoking, critical illness
- Previous or existing PU

**Other important intrinsic risk factors**
- Reduced sensation – e.g. spinal cord injury, stroke, local or general anaesthesia
- Increased skin moisture – e.g. incontinence, perspiration
- Increased age
- Anaemia
- Poor nutrition, including low blood albumin levels
- Reduced general health status – e.g. acute illness, renal impairment, carcinoma

Table 1. Examples of PU risk factors particularly relevant to risk of DTI (Berlowitz & Brienza, 2007; Black et al, 2016; Lustig et al, 2017; Preston et al, 2017; Kirkland-Kyhn et al, 2017)

<table>
<thead>
<tr>
<th>Limited mobility or immobility</th>
<th>Impaired sensation</th>
<th>Reduced tissue tolerance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Effect of risk factor</td>
<td></td>
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<tr>
<td>The patient has reduced ability to change position independently and relieve pressure</td>
<td>The patient has reduced ability to detect signals from body tissues indicating that a change in position is required to relieve pressure</td>
<td>The threshold for tissue damage induced by mechanical loading is reduced</td>
</tr>
<tr>
<td>Examples of risk factors</td>
<td></td>
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</tr>
<tr>
<td>Neurological disease, e.g. stroke, spinal cord injury</td>
<td>Prolonged sitting, e.g. in a chair, wheelchair, hard surfaces, such as toilet seats or washroom stools</td>
<td>Neuropathy, e.g. due to diabetes, alcoholism, autoimmune disease, Guillain-Barré syndrome/chronic inflammatory demyelinating polyneuropathy, malnutrition</td>
</tr>
<tr>
<td>Sedation</td>
<td>In pain, e.g. from chronic disease, recent surgery</td>
<td>Atrophy, fragility and/or breakdown of soft tissues, e.g. muscle atrophy through acute or chronic illness or paralysis</td>
</tr>
<tr>
<td>Anaesthesia, e.g. local, epidural, spinal or general</td>
<td>Prolonged confinement to a hard surface, e.g. the floor (after a fall or loss of consciousness), an emergency department trolley, radiology department surfaces, spinal board, operating theatre table or during transfer between healthcare facilities</td>
<td>Episodes of shock/hypotension/poor tissue perfusion, e.g. due to vasopressors, cardiac arrest</td>
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<tr>
<td>Unconsciousness</td>
<td>Prolonged contact with and usage of medical devices</td>
<td>Recent weight loss</td>
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<td>Malnutrition</td>
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<td>Dehydration</td>
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<td>Change in skeletal architecture, e.g. amputation, contractures, diabetic foot changes</td>
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<td>Previous PU or DTI</td>
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Diagnosis of DTI

Several conditions other than DTI can present with discoloured, purple or maroon skin, skin loss over a darkened wound bed, or rapidly developing eschar (Black et al, 2016). Healthcare professionals and carers need to be aware of the characteristics of DTI and how to differentiate it from other conditions that may present in a similar way (Table 2 below, and Appendix 1, page 30). Taking a detailed history from the patient and careful assessment will be key to establishing the correct diagnosis.

If there is uncertainty over the diagnosis of a lesion that may be a DTI, the affected area should be treated as if it is a DTI (see pages 21-26) while the concern is escalated or a referral is made to the relevant service. The urgency of the referral should be appropriate to the possible diagnosis, e.g. necrotising fasciitis is life-threatening and requires emergency referral.

Table 2. Distinguishing DTI from other conditions that present with discoloured (purple or maroon) skin +/- broken skin (Nair et al, 2014; NPUAP/EPUAP/PPPIA, 2014; Pieper, 2016; Black et al, 2016; Ousey et al, 2017b)

<table>
<thead>
<tr>
<th>Condition</th>
<th>Characteristics</th>
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| DTI (Figures 8a and 8b, page 12) | - History of application of pressure over affected area  
- Preceding event may have been 24-72 hours before recognition  
- Presents as a purple or maroon area of non-blanchable discoloured intact skin which may be surrounded by non-blanchable erythema (Box 3, page 4) +/- an overlying area of dry skin, a thin blister over a dark wound bed, a blood-filled blister, an area of broken skin with a purple, maroon, black or white wound bed, or an area of thin dark eschar (see pages 3-4).  
- Affected area may be painful, and firmer/softer, or warmer/cooler than the surrounding areas  
- Usually has a clearly defined edge  
- Most commonly affects the heels, sacrum and buttocks; may occur under a medical device |
| Category 1 PU (Figure 8c, page 12) | - History of application of pressure over affected area  
- Skin of the affected area is intact with non-blanchable reddening (erythema) or discolouration in darkly pigmented skin, and is usually over a bony prominence  
- Affected area may be painful, and firmer/softer, or warmer/cooler than the surrounding areas |
| Category 2 PU (Figure 8d, page 12) | - History of application of pressure over affected area  
- Presents as a shallow shiny or dry open ulcer due to partial thickness loss of the dermis without slough or bruising, and is usually over a bony prominence  
- May also present as an intact or open/ruptured serum-filled blister |
| Bruise (Figure 8e, page 12) | - History of blunt-force trauma is common  
- Due to extravasation of blood into tissues  
- Usually takes about two weeks to heal |
| Haematoma | - History of trauma is common  
- Due to extravasation of blood; may produce a deep-seated palpable nodule  
- Morel-Lavallée lesions are a type of haematoma that are due to trauma that separates the subcutaneous tissues from the underlying fascia and muscle; skin necrosis and infection may occur |
| Blood blister | - History of trauma is common, e.g. pinching of tissue between two hard surfaces  
- Forms a raised blister that is filled with blood; may be red initially and then become purplish or black |
| Skin tear (Figure 8f, page 12) | - History of trauma is common  
- Common on the extremities of older people  
- May cause full- or partial-thickness skin loss  
- Often accompanied by bleeding |
| Incontinence-associated dermatitis (Figure 8g, page 12) | - Usually associated with faecal and/or urinary incontinence  
- Affects perineum, peri-genital area, buttocks, gluteal fold, upper thighs and lower back  
- Affected area often has poorly defined edges  
- Erythema is blanchable and may be blotty, and may be accompanied by partial thickness skin loss and vesicles |
| Venous engorgement (Figure 8h, page 12) | - Can occur in dependent tissues in patients with poor cardiac output; may be associated with swelling due to oedema  
- The colour of the affected area can change if moved, e.g. during elevation or position change; discoloration is blanchable |
| Arterial insufficiency | - Skin may be pale or mottled red to bluish/purple colour; dry and hairless  
- Often bilateral; patient may keep the limb(s) in a dependent position and complain of numbness, burning sensation and pain  
- Slow capillary refill and pallor on elevation (Box 6, page 13); pulses in an affected limb may not be palpable  
- Embolic events: small blood clots may cause sudden onset of ischaemia with pain and discoloration of the affected tissues, e.g. in the toes |
Figure 8: DTI and other conditions that may have a similar presentation
See Table 2, page 11, for more information.
Overview of assessment, management and prevention

The principles underlying the assessment, management and prevention of DTIs are similar to those for the assessment, management and prevention of PUs (Preston et al, 2017).

The assessment, management and prevention of DTIs should be undertaken in the context of mental capacity and consent (Box 7)

FORMAL CARER, INFORMAL CARER OR PATIENT

For patients in community settings, e.g. in their own homes or in nursing homes, signs and symptoms of DTI may first be recognised by him- or herself, or by a formal or informal carer. The patient or carer noticing such signs should contact a registered healthcare professional as soon as possible, and in the meantime should:

- Protect the affected area:
  - Make sure the patient does not sit or lie on the affected area and that nothing presses on it
  - Cover broken skin or blistering with an appropriate low-adherent dressing as per local dressing formulary, if available; if not available, leave the broken or blistered area exposed and avoid touching it

- Prevent any additional areas of pressure-induced damage:
  - Continue any other PU prevention measures
  - Keep the patient clean and dry

- Encourage healthy eating and fluid intake (Figure 9, page 14).

Box 6. Clinical examination to test for arterial insufficiency

- Elevation of a limb affected by arterial insufficiency may cause pallor
- In the lower limb, Buerger’s test can be used in assessment for lower limb ischaemia (Insall et al, 1989). For example, in the lower limb, elevation of the leg to an angle of 45 degrees to the bed surface for one to two minutes when the patient is lying on his/her back will cause pallor of the skin; then when the patient sits with the leg lowered over the edge of the bed, the skin may become bluish and then reddened
- If arterial insufficiency is suspected, consider determining ABPI (ankle–brachial pressure index) and/or referral to a vascular service

Box 7. Mental capacity and consent (Department of Health, 2009; Mughal, 2014; Nichols, 2014)

- Valid consent is required before touching a patient, whether that is to assist with self-care tasks such as dressing, to examine skin as part of PU prevention, or to undertake invasive procedures
- Valid consent requires that the patient has the mental capacity to give consent, i.e. that the patient can understand and use information to make a decision
- The Mental Capacity Act 2005 covers adults aged over 16 years in England and Wales and sets out criteria for establishing mental capacity. The Act has five main principles:
  - Assume the patient has mental capacity unless it is proven otherwise by undertaking a mental capacity assessment
  - Ensure all practical steps have been taken to support the patient with the assessment before deciding they lack mental capacity
  - An unwise decision does not mean that the patient lacks mental capacity
  - Any decision made on behalf of a patient lacking mental capacity must be made in his/her best interests
  - Always consider whether there is a least restrictive option when making any best-interests decision
- A patient who has mental capacity should be provided with sufficient information in a way that they can understand to inform their decision about consent, such as the options available, the risks and benefits, and the implications of their choices
- Any decision made in relation to a patient who lacks capacity should be made in that person’s best interests
- The issues of capacity and consent are covered in Scotland by the Incapacity (Scotland) Act (2000) (Scottish Government, 2000) and in Northern Ireland by the Mental Capacity Act (Northern Ireland) 2016 (TSO, 2016)
- Healthcare professionals and carers should follow the consent policies and procedures of their organisation
The diagnosis of a DTI may be clear on presentation, but referral as appropriate, e.g. to a tissue viability service, may be needed if it is not (Table 2, page 11). Experienced healthcare professionals aware of the evolution of DTIs may decide to delay referral for a short period (e.g. up to 1–2 weeks) if the area appears stable, and the patient has no other symptoms of illness that warrant onward referral. In such cases, assessment and management should be commenced as if for a DTI alongside frequent monitoring to see whether the diagnosis becomes more evident (Figure 10, page 15).

It is important that an area showing signs and symptoms of a DTI is immediately protected from any further mechanical load even when the diagnosis is not clear; do not wait for a referral to be completed or for confirmation of the diagnosis.

Prevention of DTI should generally adopt the same principles as the management of DTI. A focus of management of DTI is the prevention of further tissue damage (Preston et al, 2017). Information on the:
- Assessment of a patient with signs and symptoms of a DTI is on pages 16-20
- Management of a patient with signs and symptoms of a DTI is on pages 21-26
- Prevention of DTI is on pages 27-28.

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**Figure 9: Overview of assessment and management of a patient with signs and symptoms of a DTI by the patient, family member or formal/informal carers, and registered healthcare professionals**

**Signs and symptoms of a DTI**
- Reported by the patient
- Detected on regular skin inspection or during routine care

**Recognised initially by:**
- Formal carer, informal carer, family member or patient
  - Refer immediately to a registered healthcare professional
  - AND without waiting for the patient to be seen, and after establishing capacity and consent (Box 7, page 13):
    - Make sure the patient does not sit or lie on the affected area, that nothing presses on it and that it is not in contact with a bed, chair, seat, device or the floor
    - Cover broken skin or blistering with a low-adherent dressing if available
    - Provide pain relief as prescribed/according to local policy
    - Keep the patient clean and dry
    - Continue using any other PU prevention measures or equipment
    - Encourage healthy eating and fluid intake

- Registered healthcare professional/allied healthcare professional
  - Diagnose DTI
  - OR
    - If the diagnosis is unclear (see Table 2, page 11), refer according to local policy for clarification, or if sufficiently experienced and the patient has no other symptoms that require referral, consider delaying referral for a short period of observation
  - AND without delay, and after establishing capacity and consent (Box 7, page 13), and in the context of a multidisciplinary team:
    - Proceed with holistic assessment (pages 16-20) and management (pages 21-26)
    - Figure 10, page 15
Figure 10: Principles of assessment, management and prevention of DTI

- Assess
  - General health
  - Nutrition and hydration
  - Skin
  - Pain
  - Level of mobility and activity
  - Continence status
  - Risk for developing further DTI/PU
  - Availability and use of offloading/pressure redistribution equipment
  - Psychosocial status
  - Insight and understanding of DTI/PUs
  - Patient/family/carer objectives
  - DTI (wound)

- Define objectives of management, e.g.:
  - Reduce risk of further tissue damage (further DTI/PU)
  - Healing of DTI
  - Reduction in pain

- Reduce risk of further tissue damage (involving the multidisciplinary team as appropriate):
  - Offloading/regular repositioning/pressure redistribution
  - Optimise patient condition - manage comorbidities and incontinence; improve nutrition and hydration; protect skin; optimise mobility
  - Manage pain, e.g. pain associated with repositioning

- Manage the DTI:
  - Offload the DTI
  - DTI wound management
  - Manage pain associated with the DTI

- Record and report DTI according to local policy for incident reporting

- Patient, family and carer education

- Monitor and reassess regularly
  - Assess DTI healing progress at least weekly; inspect skin at least daily
  - Conduct ongoing PU risk assessment
  - Effectiveness and suitability of offloading/pressure redistribution/repositioning regimen
  - Consider the patient’s other needs/management objectives for other conditions

- Refer according to local policy:
  - For clarification of diagnosis if necessary
  - If condition of the DTI is deteriorating rapidly
  - If condition of the DTI is not improving within 2 weeks (or sooner if the patient is not stable)

*Taking into account all of the patient’s needs
Assessment of DTI

A patient with a DTI should undergo a structured holistic assessment that includes:

- The patient’s overall health, including comorbidities, nutrition and skin condition
- The DTI
- Identification of the risk factors for DTI and PUs (Figure 10, page 15).

A holistic and structured assessment of a patient with a DTI will provide essential information for devising and implementing an appropriate management plan. The results of the holistic assessment must be documented according to local policy (Chamanga & Ward, 2015).

GENERAL HEALTH STATUS

Assessment of general health should include:

- Medical and surgical history – may reveal risk factors for the development of pressure-induced damage, e.g. stroke, diabetes
- History of previous PU or DTI – may indicate increased risk
- Current health – active comorbidities; new symptoms that may need investigation
- Medication – e.g. medication that causes sedation may increase risk of DTI, or chronic use of steroids may impair wound healing
- Factors that may adversely affect healing (Box 8, page 17).

Ascertainment of general health status will allow proposed management (and prevention) of the DTI to be placed in context. In some situations, e.g. in critical illness or end of life care, the other needs of the patient may take priority over DTI prevention and management. Where this is the case, such decisions should be clearly documented.

NUTRITION

Poor nutritional status and inadequate dietary intake are risk factors for PU development and impaired wound healing (NICE, 2014; Posthauer et al, 2015). Patients should be assessed for recent weight loss and problems associated with eating and drinking. The Malnutrition Universal Screening Tool (MUST) is an example of a screening tool cited by the National Institute for Health and Care Excellence (NICE, 2006; Elia, 2003).

Patients suspected of having poor nutrition should undergo nutritional assessment according to local policy, which may include referral to a dietician. Patients with difficulty eating or drinking may also benefit from assessment by an occupational therapist.

SKIN ASSESSMENT

Skin assessment should ensure examination of the patient’s entire skin for signs of pressure-induced damage, e.g. areas of non-blanchable erythema (Box 3, page 4), including under medical devices, such as face masks, and for general condition. Dry skin, particularly at the heels, is a risk factor for pressure-related damage (Lechner et al, 2017).

Checking for evidence of pressure-induced damage should involve inspecting and palpating the skin, and asking the patient about any areas of discomfort or pain: ‘look, feel, ask’

Careful attention should be paid to inspection of areas in which the patient is complaining of pain. If the area cannot be accessed easily for inspection, e.g. because of difficulties with or uncertainty over the removal of a device or bandage, the concern should be escalated as soon as possible. After removal of the device and skin inspection, careful consideration may be required of the suitability of reapplication of the device and whether there is need for an alternative.
Heels are the most common site of DTI, but may be difficult to access. Particular effort may be needed to ensure that any foot coverings (e.g. slippers, shoes, socks, tights, anti-embolic stockings) are removed regularly, e.g. at least daily or more frequently in patients at high risk, to allow examination.

Patients who are incontinent or who had been confined to the floor after falling may have signs of incontinence-associated dermatitis (Table 2, page 11).

All of the patient’s skin should be examined: patients may have more than one DTI (Figure 11), also have a PU or a skin condition that increases risk of further pressure-induced damage

Skin assessment provides an opportunity to talk to the patient about risk for DTIs and to provide advice on avoidance of pressure-induced damage and the signs and symptoms if it does occur.

PAIN ASSESSMENT
Assessment of pain is important because:

- DTIs can cause pain
- Pain may occur before physical signs of pressure-related damage occur
- In a DTI with an open wound, a sudden onset or an increase in pain may indicate infection
- Pain and psychological stress can adversely affect wound healing
- Pain occurring during movement may reduce mobility, compounding risk for further pressure-induced damage (Gorecki et al, 2009; Woo, 2012; Black et al, 2016).

Repeated use of the same method of assessment will assist in monitoring pain and the effectiveness of pain-relieving interventions. The method of pain assessment should be selected according to the patient’s cognitive and linguistic abilities. In patients with reduced cognitive ability, non-verbal cues, body language and the family/carers’ knowledge of the patient may aid assessment of pain (RNAO, 2016).

Patients should be assessed for DTI-related pain and non-DTI-related pain. Assessment should include evaluating the location, nature, frequency, timing, and severity of pain (Box 9) and factors that initiate or relieve pain

Box 8. General factors that may impair wound healing (Guo & DiPietro, 2010; Anderson & Hamm, 2012)

- Extremes of age
- Diabetes
- Peripheral arterial disease
- Severe liver or kidney disease
- Obesity
- Immunosuppression – e.g. radiation therapy, AIDS
- Medication – e.g. steroids, immunosuppressants, chemotherapy, non-steroidal anti-inflammatory drugs (NSAIDs)
- Smoking
- Alcoholism
- Stress
- Malnutrition

Box 9. Examples of scales for assessment of pain severity (Solowiej & Upton, 2010)

- **Verbal pain rating scale** – the patient indicates from a list of phrases ranging from ‘no pain’ to ‘severe pain’ which phrase describes their pain best
- **Numerical pain rating scale** – the patient is asked to score their pain on a 0 to 10 scale, where 0 = no pain and 10 = the worst pain imaginable
- **Visual analogue scale** – the patient is presented with a line, one end of which represents no pain and the other end the worst pain imaginable, and is asked to draw a mark on the line to represent the level of pain
LEVEL OF MOBILITY AND ACTIVITY

A central principle of the management (and prevention) of DTI is removal of any mechanical load from the affected area and any other areas at risk of pressure-induced damage. A patient who is able to perform purposeful movements, i.e. is able to lift or tilt a body part completely away from the surface they are resting on, can be encouraged to change position regularly as part of a pressure-relieving regimen.

RISK FOR FURTHER PRESSURE-INDUCED DAMAGE (DTI OR PU)

A patient with a DTI is at risk of developing a further DTI or a PU. An understanding of the risk factors for DTI and PUs (Box 5 and Table 1, page 10) will enable healthcare professionals, carers, patients and family members to quickly identify which risk factors are relevant to the patient. This should include understanding that development of an acute illness in a patient already at risk of pressure-related damage will further increase risk. Consideration should also be given to all aspects of care that may be associated with risk, e.g. prolonged sitting without position change in a chair or on a toilet (Lustig et al, 2017).

The patient’s level of mobility, along with identification of DTI or PU risk factors and aspects of care that may be associated with risk for pressure-induced damage can be used to inform the management plan.

PU risk assessment tools for different patient groups are available and should be used according to local policy (Table 3). Currently, there is no risk assessment tool specific to DTIs.

SUPPORT SURFACES AND PRESSURE REDISTRIBUTION EQUIPMENT

All support surfaces and pressure redistribution equipment in use by the patient, including its physical condition, suitability and whether use is appropriate and safe, should be assessed (RNAO, 2016). Examples of equipment to consider include bed, chair/wheelchair, toilet seat, bathing stool, foot rest, transfer equipment, and interventional medical device.

OTHER

- Psychosocial status – including for patients in the community, the formal and informal care systems that may be in place.
- Understanding of DTI and PUs – ascertaining the level of insight into DTI and PUs of the patient, family, and, where appropriate, the carer may reveal the need for education to aid understanding of management strategies and involvement in care planning.

Table 3. PU risk assessment tools (Wounds UK, 2013a; Fletcher, 2017)

<table>
<thead>
<tr>
<th>Patient population</th>
<th>Appropriate risk assessment tool</th>
</tr>
</thead>
</table>
| Adults             | Acute care: Waterlow Score (Waterlow, 2005); PURPOSE T (Coleman et al, 2017)  
|                    | Critical care: Jackson/Cubbin Pressure Area Risk Calculator (Jackson, 1999)  
|                    | Community care: Walsall Community Pressure Sore Risk Calculator (Chaloner & Franks, 2000)  
|                    | Orthopaedic: Pressure Sore Prevention Score (Lowthian, 1989) |
| Older people       | Norton Pressure Ulcer Risk-Assessment Scale (Norton, 1989)  
|                    | Braden Scale for Predicting Pressure Sore Risk (Bergstrom & Braden, 1992) |
| Paediatric patients| Glamorgan Paediatric Pressure Ulcer Risk Assessment Scale (Willock et al, 2009)  
|                    | Braden Q Scale (Curley et al, 2003) |
DTI ASSESSMENT
Assessment of a DTI will provide a baseline from which to monitor progress or detect deterioration. The assessment should be clearly documented according to local policy. Serial photographs may be helpful and should be obtained after gaining patient consent and stored according to local policy (Farid et al., 2014; Sperring & Baker, 2014).

The assessment should include, as a minimum, the location of the DTI and measurement of the size of the affected area and blistering or open wound if present. Systematic assessment of the DTI will be aided by using a wound assessment framework (Box 10), even when the skin over the DTI remains intact.

If the skin is broken, the wound bed should be assessed for the proportions of granulation tissue, necrotic tissue/eschar, slough and epithelial tissue present (Ousey & Cook, 2012). Exudate levels, though difficult to assess, influence dressing selection. Exudate consistency, colour and smell may also indicate infection (Wounds UK, 2013b). See Box 11 for a summary of signs of possible local wound infection. If bone can be probed in the base of the wound or is exposed, osteomyelitis should be suspected (Rennert et al., 2009). Systemic signs of wound infection include malaise and pyrexia.

In patients with more than one DTI, or a DTI and another wound or area of skin damage such as incontinence-associated dermatitis, each area of damage or wound should be individually assessed and categorised to facilitate appropriate care planning and management.

Box 10. Frameworks to aid systematic wound assessment
- TIME(S) (Schultz et al., 2004; Wounds UK, 2016)
  - Tissue
  - Infection/inflammation
  - Moisture imbalance
  - Edge of the wound
  - Surrounding skin
- Triangle of Wound Assessment (Dowsett et al., 2015)
  - Wound bed
  - Wound edge
  - Periwound skin

Box 11. Signs and symptoms of possible wound infection in a DTI with broken skin or an open wound (WUWHS, 2008)
- New, increased, or altered pain
- Malodour or change in odour
- Increased or altered/purulent exudate
- Delayed healing
- Periwound oedema
- Bleeding or easily damaged granulation tissue
- Altered wound bed colour
- Induration of periwound skin
- Pocketing and bridging
The Expert Working Group recommends that a DTI remains recorded as a DTI, and is not recategorised at any point (e.g. to a Category 3 or 4 PU), even if it evolves.

The Expert Working Group recommends that the notes of a patient with a DTI are flagged to alert all healthcare professionals because of the likely increase in risk for further pressure-induced damage.

**DIAGNOSTIC TESTS FOR DTI**

The difficulties of identifying the early stages of DTI have generated interest in the potential of tests that may indicate the presence of tissue damage. Tests under investigation for the detection of pressure-induced damage include assessing biomarkers in plasma, sweat and urine related to muscle damage and changes in concentrations of inflammatory mediators (Ferguson-Pell & Hagisawa, 1988; Knight et al, 2001; Loerakker et al, 2012; de Wert et al, 2015; Krishnan et al, 2016; Worsley et al, 2016).

Ultrasound scanning (USS) can detect subcutaneous tissue changes in DTI prior to changes becoming visible on the skin surface (Scheiner et al, 2017). Therefore, USS has potential as a screening tool in patients considered to be at risk. However, the clinical utility of USS for this indication is likely to be limited because of the resource, training, time and cost implications of widening access to USS.

In areas of pressure-induced damage, the amount of interstitial fluid increases in response to the inflammatory process. As a result, hand-held devices to measure subepidermal moisture (SEM) have been developed to aid early identification of PUs (Moore et al, 2016). Clinical studies have reported that SEM values increase with increasing tissue damage (Oliveira et al, 2017).

Thermography has been investigated to assess its ability to predict progression of discoloured intact skin to necrosis (Cox et al, 2016). The study found that intact discoloured areas with cooler skin temperatures in the centre in comparison with surrounding skin were significantly more likely to develop necrosis by day 7 than if the skin was warmer than the surrounding areas. However, nurses participating in the study were unsure of the feasibility of using thermography in clinical practice and more investigation is required.

**Currently, there is no widely accessible and reliable test to aid diagnosis of DTI**

**A DTI may be subject to reporting and investigation according to local policy**
Management of DTI

The management and prevention of a DTI share many similarities. In fact, prevention strategies are used in DTI management to aid healing and prevent deterioration of the DTI and to prevent any further pressure-induced damage (Figure 12). Management of DTI therefore comprises local treatment of the DTI plus DTI prevention. Prevention of DTI is discussed on pages 27-28.

The assessment of the patient with a DTI will indicate appropriate objectives that will form the basis of the management plan. The plan needs to take into consideration all the patient's needs. In some cases, the DTI aspects of the plan may need to be adjusted or take lower priority.

The objectives and plan should be discussed and agreed where possible with the patient, family and carers. Management of DTI is likely to require a multidisciplinary team approach, which may also include as appropriate nurses, tissue viability nurses, healthcare assistants, physiotherapists, occupational therapists, podiatrists, dieticians, medical specialists, surgeons, bioengineers and social workers (RNAO, 2016).

Patients with a DTI often have other conditions. In some situations, e.g. in patients who are haemodynamically unstable, the management of these conditions needs to take priority over, or may complicate, the management (and prevention) of DTIs or PUs. In such cases, this may mean that it is necessary to recognise that the DTI is likely to evolve and deteriorate.

OPTIMISE PATIENT CONDITION AND NUTRITION

Optimising the patient’s condition aims to remove or ameliorate any modifiable risk factors for pressure-induced damage, e.g. comorbidities or incontinence, and any factors that may impede healing identified during assessment.

Patients should be encouraged to eat healthily and to maintain good levels of hydration. If a patient is unable to achieve an adequate nutritional intake with meals, fortified foods and/or high protein oral nutritional supplements or enteral/parenteral nutritional support as appropriate should be considered (NPUAP/EPUAP/PPPIA, 2014).

OFFLOADING, PRESSURE REDISTRIBUTION AND REPOSITIONING

As mechanical loading, mainly in the form of pressure, is the main cause of DTI, it is logical to remove pressure from the affected area and, as far as possible, reduce the intensity and duration of pressure over other areas that may be at risk of pressure-induced damage.

Wherever possible, it is important to offload a DTI and avoid exposing it to any form of mechanical load.

In an at-risk patient who reports pain in an area in which a DTI may occur, but who has no other signs of DTI, the affected area should be offloaded if possible and monitored very frequently.

A reduction in mechanical load can be:

- Partial – i.e. produce a reduction in mechanical load, known as pressure redistribution; this may be intermittent or continuous.
- Complete – i.e. totally remove mechanical load, known as offloading; this may be intermittent or continuous.

In practice, regular repositioning, which is used to offload areas, and equipment, which may redistribute pressure or offload, are often used in combination. Table 4, page 22, provides information on offloading according to the most common locations of DTI.
### Table 4. Offloading for a DTI according to location (Levy et al, 2015; WUWHS, 2016; Stephens & Bartley, 2017)

<table>
<thead>
<tr>
<th>Location of DTI</th>
<th>Tips on repositioning/offloading/pressure redistribution</th>
</tr>
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<tbody>
<tr>
<td>Heel</td>
<td>■ Reposition the patient to offload pressure from the heels when sitting or lying by ensuring they are not in contact with the bed, floor, foot rest etc, i.e. the heels are ‘suspended’ or ‘floating’&lt;br&gt;■ Devices that can be used to aid heel offloading:&lt;br&gt;  - Boot – air, gel, foam, fibre&lt;br&gt;  - Heel zone mattress&lt;br&gt;  - Pillow or cushion (Figures 13 and 14, pages 24 and 25)&lt;br&gt;  - Silicone cup&lt;br&gt;  - Trough&lt;br&gt;  - Wedge&lt;br&gt;■ Rigid framed devices may be needed for ambulant patients&lt;br&gt;■ On intact skin, consider use of a dressing designed to reduce exposure of the DTI to mechanical loads* (see page 26 for the management of DTI with broken skin or an open wound)</td>
</tr>
<tr>
<td>Sacrum/ischial tuberosity</td>
<td>■ Reposition the patient to offload the sacrum/ischial tuberosities when lying or ask the patient to reposition themselves if possible and safe to do so&lt;br&gt;■ In bed, use the 30-degree tilt side-lying position (Figure 14, page 25) and avoid the patient lying on their back (supine)&lt;br&gt;■ Minimise time seated; avoid sitting patients with an ischial DTI in a fully erect posture; patients should be encouraged to offload intermittently while sitting, e.g. by lifting themselves completely off the seat, sideways leans, forward tilts and leans, and standing&lt;br&gt;■ On intact skin, consider use of a dressing designed to reduce exposure of the DTI to mechanical loads* (see page 26 for the management of DTI with broken skin or an open wound)</td>
</tr>
<tr>
<td>Under a medical device</td>
<td>■ Where possible, change the location or shape of the medical device, e.g. for oxygen masks, use a different mask shape/size and/or alternate mask use with nasal cannulae&lt;br&gt;■ Regularly lift and reposition the device (at least as frequently as general repositioning), ensuring that the tension of any holding straps or fixation is sufficient for function of the device but is not higher than necessary&lt;br&gt;■ The use, positioning and repositioning of medical devices in patients who are oedematous requires careful consideration&lt;br&gt;■ On intact skin, consider use of a dressing (see page 26 for the management of DTI with broken skin or an open wound); care should be taken not to select a dressing that does not interfere with the function of the device and that does not increase pressure, e.g. is not too thick</td>
</tr>
</tbody>
</table>

*Dressings with low friction outer surfaces may reduce superficial mechanical loads. Dressings with multiple layers have been found in computer models of the heel to dissipate mechanical loads at the bone:tissue interface more effectively than single-layer dressings; this effect may be due in part at least to the horizontal displacement of the different dressing layers relative to each other and to a cushioning effect.

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Even when equipment for offloading and pressure redistribution is in use, the patient should be repositioned regularly, unless contraindicated, and should undergo regular reassessments and skin inspections (NPUAP/EPUAP/PPPIA, 2014)

Patient preference, comfort and concordance should be taken into account when offloading and repositioning

**Repositioning**

Frequent repositioning is an important method of intermittently offloading the tissues.

Positioning can also be used to avoid applying pressure to an existing DTI. The exact nature of the repositioning regimen will depend on the:

■ Patient’s level of mobility and whether, and to what extent, they can reposition themselves<br>■ Patient’s ability to tolerate the new position<br>■ Location of existing pressure-induced damage – e.g. while in bed a patient with a sacral DTI might be positioned on their left and right sides only and not on the back<br>■ Location of the patient – e.g. in bed or a chair<br>■ Support surface in use (NPUAP/EPUAP/PPPIA, 2014).
A repositioning regimen should describe the positions to be used along with the frequency and duration of the position change, and should be reviewed regularly (Box 12) (NPUAP/EPUAP/PPPIA, 2014). The frequency of repositioning will depend on patient risk. Those who are at higher risk and are more susceptible to pressure-induced damage need to be repositioned more frequently.

As a guide, patients should be repositioned every 2–4 hours when lying on a pressure redistributing mattress and should shift their weight every 15 minutes if sitting (NPUAP/EPUAP/PPPIA, 2014; RNAO, 2016). Any aids used to assist with repositioning (e.g. hoists, glide sheets) should prevent further tissue trauma and use should be documented. After repositioning, sheets, garments and incontinence products should be wrinkle-free. In general, the number of layers between a patient’s skin and a therapy surface should be kept to a minimum.

Devices including pillows, wedges, gel or fluidised mouldable positioners can be used to support a patient in a new position (Preston et al, 2017; NPUAP/EPUAP/PPPIA, 2014). Beds that allow the angle of the head and knee/foot sections to be adjusted independently can aid patient positioning (Preston et al, 2017).

Offloading and repositioning do not necessarily require specialist equipment; pillows can be used to good effect to support a patient in a position or can be used to provide offloading, e.g. of the heels if placed under the lower legs, or of bony prominences if placed either side.

Independent movement by a patient can move a body part away from a preventative device, e.g. if the legs are moved from pillows used for heel offloading, the heels may come into contact with the support surface. Consider the use of an appropriate alternative, e.g. a patient-worn removable or non-removable device or a mattress with inbuilt heel zone, if independent movement is problematic.


- Avoid whenever possible positioning a patient directly on to a DTI (or a PU)
- Where safe and possible encourage patients to reposition themselves by lifting vulnerable tissue areas clear of the support surface
- When assisting or moving a patient to a new position:
  - Follow local manual handling procedures and policies
  - Avoid ‘dragging’ the patient along the support surface as this creates external shear forces
- Ensure manual handling equipment and medical devices are not left under the patient, unless designed to be left in place
- Repositioning in a bed:
  - Use the 30-degree tilted side-lying position (Figure 14, page 25) and alternate moving from right side to back to left side as appropriate; if this cannot be tolerated use the prone position if not contraindicated
  - Avoid 90-degree side-lying or semi-recumbent postures
  - Limit head of bed elevation to 30 degrees or less, unless medically contraindicated or if higher elevation is required for feeding, digestive or respiratory considerations
- Pillows, wedges, and gel or fluidised positioners can aid positioning
  - If necessary, a knee break (gatch) can be used to reduce the risk of the patient sliding down the bed
  - For taller patients, consider use of bed extensions and foam squabs to avoid feet being wedged against the foot plate of the bed
  - Reposition every 2–4 hours or more frequently if there are signs of deterioration
- Repositioning while seated (in a chair or wheelchair)
  - Ensure the chair is suitable for the size and weight of the patient and fits correctly (Box 13, page 24)
  - Select a seating position that:
    - Is stable
    - Is acceptable to the patient and allows them to carry out their full range of functions
    - Ensures support for the feet (on the floor or appropriate footrest) and arms
  - Reassess the effectiveness of the repositioning regimen regularly with respect to enabling the DTI to heal and prevention of other areas of pressure-induced damage

After repositioning, always check that the area(s) of concern are properly offloaded
Pressure redistribution
Patients with a DTI, or at increased risk of a DTI or other pressure-induced damage, should be placed on a pressure redistribution support surface selected according to local policy. Pressure redistribution support surfaces distribute the mechanical load more uniformly and so reduce the load or pressure over bony prominences (International Review, 2010). Pressure redistribution support surfaces are available in a wide range of formats including mattresses, integrated bed systems, mattress overlays, seat cushions and toilet seats.

Support surface selection should consider the patient’s needs and should be appropriate for use in the care setting (NPUAP/EPUAP/PPPIA, 2014)
Pressure redistribution surfaces should be considered for the bed and seating (including chairs, wheelchairs, toilet seats and bathing/showering stools) (Table 5, page 25). Air-fluidised and alternating pressure surfaces can be useful for patients who are acutely ill and/or who cannot be repositioned (Ovens, 2012).

Support surfaces should not replace good care and repositioning, and their use should be reassessed regularly
As the patient improves, the support surfaces in use should continue to be reassessed carefully to ensure suitability and that the surface does not hamper independent movement.

PAIN MANAGEMENT
In addition to management of any background pain, pain management may need to be considered for repositioning and during dressing changes or debridement.
SKIN INSPECTION AND MANAGEMENT
The patient’s skin should be inspected regularly (at least daily) for signs of pressure-induced damage including non-blanchable erythema and DTI (NPUAP/EPUAP/PPPIA, 2014). The skin under medical devices should be inspected at least twice daily for signs of pressure-induced damage (NPUAP/EPUAP/PPPIA, 2014).

Regular skin inspection is an essential part of DTI management and of pressure-induced damage prevention

The patient’s skin should be kept clean and dry. Cleansing should use a pH-balanced skin cleanser. Dry skin can be moisturised with a non-sensitising, fragrance and alcohol-free moisturiser (Norton et al, 2017).

If the patient is incontinent, reversible causes such as urinary tract infection or constipation should be treated. The skin should be cleansed at least once daily or after each episode of faecal incontinence. A skin protectant/barrier should be applied to areas with or at risk of incontinence-associated dermatitis (Beeckman et al, 2015; Fletcher, 2015).

<table>
<thead>
<tr>
<th>Table 5. Types of pressure redistributing support surfaces (Ovens, 2012; Fletcher et al, 2015)</th>
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<tbody>
<tr>
<td>Type of support surface</td>
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<tr>
<td>------------------------</td>
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<tr>
<td>Reactive support surfaces</td>
</tr>
<tr>
<td>Foam</td>
</tr>
<tr>
<td>Air- or gel-filled</td>
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<tr>
<td>Low air loss</td>
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<tr>
<td>Air-fluidised</td>
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<tr>
<td>Active support surfaces</td>
</tr>
<tr>
<td>Alternating pressure</td>
</tr>
<tr>
<td>Hybrid support surfaces</td>
</tr>
<tr>
<td>Non-powered</td>
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<tr>
<td>Powered</td>
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</table>

LOCAL DTI MANAGEMENT
The approach to the local management of a DTI depends on whether the skin is intact.

**DTI with unbroken skin**
A DTI with unbroken skin can be left uncovered if acceptable to the patient, family and carers. If a dressing is applied, it should have a low friction outer surface. The skin under the dressing should be inspected at least once daily and the dressing changed in accordance with the manufacturer’s instructions (WUWHS, 2016). In general, blisters should be left intact. On occasion, a blister may be deroofed under aseptic conditions, e.g. if a heel blister is preventing mobilisation.
**DTI with broken skin or an open wound**

A DTI with broken skin or an open wound should be cleansed and covered with a dressing that is appropriate for exudate level, wound bed tissue type and condition of the periwound skin with the aim of promoting moist wound healing (NPUAP/EPUAP/PPPIA, 2014). The cleansing agent and dressing should be selected according to local policy. If exudate levels are high, negative pressure wound therapy may be appropriate (Wounds UK, 2013b).

In general, debridement of a DTI should be approached conservatively. Surgical debridement is reserved for severe cases. An antimicrobial dressing can be considered for an ‘open’ DTI that is at high risk of infection or that is showing clinical signs and symptoms of infection (Box 11, page 19) (NPUAP/EPUAP/PPPIA, 2014).

The choice of dressing applied to a DTI with broken skin or an open wound should be based on the needs of the wound as determined by structured wound assessment and should follow local dressing formulary guidance.

Healing progress should be monitored at least weekly and documented. For a DTI with an open wound, a tool such as the Bates-Jensen Wound Assessment Tool (BJWAT), the Pressure Ulcer Scale for Healing (PUSH) or the Pressure Sore Status Tool (PSST) may aid monitoring of healing progress (NPUAP/EPUAP/PPPIA, 2014).

**PATIENT, CARER AND FAMILY EDUCATION**

Education should include what a DTI is, how it is caused, how to recognise it, what the treatment is, how it can be prevented and where to get help or further information. Management of expectations including time frames and likely outcomes, where possible, should also be included.

The content and mode of delivery should be tailored to the needs of the individual. A combination of educational methods may be helpful. A discussion allows the information presented to be tailored to the understanding of the individual and the opportunity to ask questions.

Combining discussions with other methods of providing information, e.g. printed leaflets and website links, will give further opportunities for learning and reflection (Colledge et al, 2008).

**MONITORING AND REASSESSMENT**

The patient should receive ongoing risk assessment and should be reassessed if there is a change in his/her condition. The effectiveness and suitability of the offloading, pressure redistribution and repositioning regimen should be reviewed.

Referral to a tissue viability service may be needed if the condition of the DTI:
- Deteriorates rapidly or
- Does not improve within two weeks or within a timeline defined in local policies.

**DOCUMENTATION**

All aspects of the patient’s management plan should be documented, along with rationale for each element and timing for review. The patient’s notes should be flagged to show that they have had a DTI because of the likely increase of increased risk for future pressure-induced damage.

The presence of a DTI, along with current management, should be clearly communicated to the receiving organisation or department if a patient is transferred or admitted elsewhere.
Prevention of DTI

Guidance from NICE does not specifically address assessment of risk for DTI, but states that an assessment of PU risk should be carried out and documented for adults:

- "... being admitted to secondary care or care homes in which NHS care is provided"
- "... receiving NHS care in other settings (such as primary and community care and emergency departments) if they have a risk factor" (NICE, 2014).

The 2005 NICE guidelines state that the initial assessment should be conducted within six hours of the first episode of care in a setting (NICE, 2005).

Healthcare Improvement Scotland’s standards for prevention and management of PUs states that PU risk should be assessed and documented:

- “Within 8 hours of admission to hospital or care home
- Within 24 hours of admission to any other care setting
- On the first visit from community services or teams, for example, community nurse, hospital at home, social care or care at home” (HIS, 2016).

The NPUAP/EPUAP/PPPIA guidelines recommend that a structured risk assessment is carried out “as soon as possible (but within a maximum of eight hours after admission)” (NPUAP/EPUAP/PPPIA, 2014).

ASSESSMENT OF RISK FOR DTI

Patients should be reassessed regularly and according to local policy for risk for pressure-induced damage. Reassessment of risk should also be carried out if the patient’s condition changes and/or they develop an additional risk factor (Box 5 and Table 1, page 10) (NPUAP/EPUAP/PPPIA, 2014).

Patients, families and carers should be alert to any indicators or changes that may indicate increased risk for DTI or pressure-induced damage and contact a healthcare professional for consideration of a formal assessment.

Underpinning DTI prevention are:

- Formal assessment of risk for pressure-induced damage
- Regular skin inspections to detect early signs of pressure-induced damage, e.g. non-blanchable erythema.

PU risk assessment tools

PU risk assessment tools for different patient groups are listed in Table 3 (page 18) and should be used according to local policy. Currently, there is no risk assessment tool specific to risk for DTI.

PU risk assessment tools provide a useful indication of level of risk for DTI and PUs. However, clinical judgement that considers all risk factors also has a very important and complementary role in guiding risk management (NPUAP/EPUAP/PPPIA, 2014).

Regular ‘head-to-toe’ inspection of the patient’s skin is an essential element of DTI prevention and should not be replaced by a PU risk assessment tool.
ASSESSMENT AND MANAGEMENT OF A PATIENT AT RISK OF DTI

The assessment and management of a patient considered at increased risk of DTI needs to take account of mental capacity and consent, patient preferences and the patient’s other needs. Figure 10, page 15, outlines the assessment and management of a patient considered at increased risk of DTI:

- Holistic assessment (see pages 16-20)
- Take action to reduce risk of tissue damage (see pages 21-26), including:
  - Offload, where possible, the anatomical locations at risk of DTI
  - Regular repositioning and pressure redistribution
  - Optimise patient condition, including:
    • Nutrition
    • Pain management
    • Management of comorbidities
- Patient, family and carer education (see page 26)
- Monitoring and reassessment, including (see page 26):
  - Regular skin inspections
  - Repeated PU risk assessments
  - Evaluation of the effectiveness of the offloading/pressure redistribution/repositioning regimen.

Patients being transferred from one facility to another may be at increased risk of DTI. The level of risk for a patient should be communicated clearly to the receiving facility. Appropriate pressure-related damage prevention measures and support surfaces should be implemented during transfer.

If signs of pressure-induced damage are detected, e.g. non-blanchable erythema, the patient and the DTI/PU prevention strategies in place should be reviewed and revised as appropriate.
Research needs and health economics

Further research is needed to establish many aspects of DTI, including incidence and the costs of management and prevention (Box 14).

Little is known about the health economics of DTI in the UK. However, from what is known of the incidence and costs of managing PUs, the socioeconomic impact of DTIs is likely to be considerable (Box 15).

Box 14. Research needs

- Large-scale studies of the epidemiology* of DTI in the UK to answer questions such as:
  - What is the incidence of DTI, including:
    • At different anatomical locations?
    • In different healthcare settings?
  - What factors are associated with increased risk of DTIs and how do they differ from those for PUs?
  - Are some risk factors for DTI more important than others?
  - What proportion of DTIs:
    • Resolve before the skin breaks?
    • Evolve to become open wounds?
  - Are there predictors of resolution and evolution of DTIs?
- Assessment of the quality of life impact of DTI
- Health economic assessment of the prevention and management of DTIs

*The most efficient way to collect large-scale data is by using data in existing registries provided that reporting is comprehensive, and includes, for example, pressure-induced damage related to medical devices. In the UK, PUs and DTIs are recorded in incident reporting systems. However, currently, there is variation in when and how DTI is recorded. In some places, DTI is not reported in monitoring systems, while in others it is observed and only recorded when it can be assigned a pressure ulcer category (1 to 4) (Coleman et al, 2016). In other places, a DTI that evolves is re-categorised as a Category 3 or 4 PU.

To aid consistency and data analysis, the Expert Working Group proposes that a DTI is recorded when first observed, and that it remains categorised as a DTI even if it resolves or evolves further with skin breakdown.

Box 15. Incidence and costs of PUs

- Incidence of PUs has been estimated to be 153,000 per year in the UK (Guest et al, 2015)
- Cost of healing a PU increases significantly with severity (from £1214 for a Category 1 PU to £14,108 for a Category 4 PU) (Dealey et al, 2012)
- Using 2013/2014 prices, the cost of managing PUs in the UK was estimated to be £507–£531 billion per year (Guest et al, 2017)

Recommended reading


*Available at: www.woundsinternational.com
**Available at: www.wounds-uk.com
### Appendix 1

**DISTINGUISHING DTI FROM RARE CONDITIONS THAT PRESENT WITH DISCOLOURED (PURPLE) SKIN +/- BROKEN SKIN** (Mustafa et al, 2009; Harr & French, 2010; Misiakos et al, 2014; Gameiro et al, 2015; Vaiman et al, 2015; Black et al, 2016)

<table>
<thead>
<tr>
<th>Condition</th>
<th>Characteristics</th>
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<tbody>
<tr>
<td><strong>Calciphylaxis</strong></td>
<td>■ May be excruciatingly tender and produce skin nodules or plaques; in later stages areas of painful necrosis produce non-healing ulcers with deep, black eschar that are often followed by infection and gangrene&lt;br&gt;■ Associated with chronic renal failure; rare in the general population&lt;br&gt;■ Due to vascular calcification and skin necrosis&lt;br&gt;■ Most commonly seen in the lower extremities and at adipose tissue sites rather than over bony prominences</td>
</tr>
<tr>
<td><strong>Perirectal abscess</strong></td>
<td>■ Presents with a dull aching or throbbing pain in the perineal area; pain worsens with sitting and on defaecation but eases after defaecation&lt;br&gt;■ May produce a tender, fluctuant mass at the anal margin</td>
</tr>
<tr>
<td><strong>Necrotising fasciitis</strong></td>
<td>■ Rare, severe, potentially lethal infection of the skin, soft tissues and muscles that may involve the abdominal wall, extremities or perineum/genitalia (Fournier’s gangrene)&lt;br&gt;■ Patients report severe pain and sometimes a history of trauma, and rapidly become unwell with sepsis&lt;br&gt;■ Skin may look pale initially, then red or bronze, and become warm and swollen. In later stages, the skin turns violet and develops large blisters and areas of gangrene&lt;br&gt;■ Requires rapid treatment with antibiotics and surgical debridement</td>
</tr>
<tr>
<td><strong>Warfarin necrosis</strong></td>
<td>■ Usually occurs within a few days of starting warfarin treatment&lt;br&gt;■ Patients are often premenopausal women being treated for deep vein thrombosis or pulmonary embolism&lt;br&gt;■ Affects the buttocks and thighs with paraesthesia and flushing of the skin followed by a <em>peau d’orange</em> appearance; haemorrhagic blisters develop within 24 hours</td>
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<tr>
<td><strong>Ecthyma gangrenosum</strong></td>
<td>■ Uncommon skin infection that may be caused by bacteria such as <em>Pseudomonas aeruginosa</em> or by fungi; reported in immunocompromised and healthy individuals; patients may develop sepsis&lt;br&gt;■ Starts with an erythematous area that develops a nodule or haemorrhagic vesicles and progresses to a necrotic ulcer, often containing eschar</td>
</tr>
<tr>
<td><strong>Gluteal compartment syndrome</strong></td>
<td>■ Rare; due to necrosis of the gluteal muscles following disruption of the artery (hypogastric) supplying blood to the muscles due to prolonged immobilisation, drug abuse, alcohol intoxication or surgery&lt;br&gt;■ Causes severe buttock pain at rest and on movement of the hip, bruising, altered sensation and swelling of the buttock&lt;br&gt;■ Usually requires surgical treatment</td>
</tr>
<tr>
<td><strong>Pyoderma gangrenosum</strong></td>
<td>■ Rare; occurs in patients with autoimmune or neoplastic diseases and most often in middle-aged adults&lt;br&gt;■ Lesions form tender papules, vesicles or pustules that evolve into painful ulcers that enlarge rapidly over a few days and that have raised violet-coloured (violaceous) and undermined borders</td>
</tr>
<tr>
<td><strong>Toxic epidermal necrolysis/ Stevens-Johnson Syndrome</strong></td>
<td>■ Rare; potentially fatal&lt;br&gt;■ Causes haemorrhagic erosions, erythema, blisters and areas of denuded skin&lt;br&gt;■ Usually due to an adverse reaction to medication, but may be due to infection with <em>Mycoplasma pneumoniae</em> or <em>Herpes simplex</em></td>
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</table>
References


