Surgical wound dehiscence
Improving prevention and outcomes
Surgical wound dehiscence (SWD) is a significant issue that affects large numbers of patients and is almost certainly under-reported. The impact of SWD can be considerable: increased mortality, delayed hospital discharge, readmission, further surgery, delayed adjuvant treatment, suboptimal aesthetic outcome and impaired psychosocial wellbeing.

Consequently, it is imperative to raise awareness of SWD and improve identification, prevention and management. Prevention of SWD comprises excellence in surgical practice, prevention of surgical site infection, reducing risk of healing impairment and use of strategies such as single-use negative pressure wound therapy in appropriate high-risk patients. Management also involves a holistic approach that includes amelioration of impediments to healing, optimising conditions in the wound bed and using appropriate treatment modalities to ultimately close the wound.

The need for international consensus on the core issues around SWD arose from the doctoral research of Kylie Sandy-Hodgetts. The process started with a meeting of an international group of surgical care experts in July 2017. Development of the subsequent consensus document included extensive review by the Core Expert Working Group and a Review Panel.

This consensus document is aimed at clinicians in all care settings who work with patients with surgical incisions. The main objective of the document is to inspire clinicians to improve outcomes for patients by providing practical guidance on how to improve prevention and management of SWD.

Karen Ousey
Chair, Core Expert Working Group

Core Expert Working Group
Karen Ousey (Chair), Professor of Skin Integrity, School of Human and Health Sciences, Director Institute of Skin Integrity and Infection Prevention, University of Huddersfield, UK; Adjunct Clinical Professor, Queensland University of Technology, Australia
Risal Djohan, Vice Chairman and Microsurgery Fellowship Program Director, Department of Plastic Surgery, Co-Director Regional ASC Quality Improvement Officer, Cleveland Clinic, Cleveland, Ohio, USA
Caroline Dowsett, Clinical Nurse Specialist, Tissue Viability, East London NHS Foundation Trust, London, UK; Independent Tissue Viability Nurse Consultant, UK
Fernando Ferreira, General Surgery Consultant, Department of Surgery: Upper Gastrointestinal and Abdominal Wall Surgery, Pedro Hispano Hospital, Matosinhos-Porto; Department of General Surgery, CLUF Porto Hospital, Matosinhos-Porto, Portugal
Theresa Hurd, Professor, Graduate Nursing, New York, USA; Clinical Nurse Specialist/Nurse Practitioner, President, Nursing Practice Solutions, Ontario, Canada
Marco Romanelli, Professor, Dermatology Unit, Department of Clinical and Experimental Medicine, University of Pisa, Italy
Kylie Sandy-Hodgetts, Clinical Trials Coordinator, Ramsay Health Care, Joondalup Health Campus, Perth; Adjunct Research Fellow, School of Human Sciences, University of Western Australia, Perth, Australia; PhD Scholar

Review Panel
Fiona Downie, Nurse Consultant - Tissue Viability, Papworth Hospital NHS Foundation Trust, UK
Franck Duteille, Professor and Head, Plastic, Reconstructive and Aesthetic Surgery, Burn Centre, University Hospital of Nantes, Nantes, France
Caroline Fife, Professor of Geriatrics, Baylor College of Medicine, Houston, Texas, USA; Medical Director, CHI St. Luke’s Wound Clinic, The Woodlands Hospital, Houston, Texas, USA; Chief Medical Officer, Intellicure, Texas, USA
Rei Ogawa, Professor and Chief, Department of Plastic, Reconstructive and Aesthetic Surgery, Nippon Medical School, Tokyo, Japan
Heidi Sandoz, Tissue Viability Services Lead, Hertfordshire Community NHS Trust, Hertfordshire, UK
James Stannard, Chair, Department of Orthopaedic Surgery; Medical Director, Missouri Orthopaedic Institute; Hanjörg Wyss Distinguished Professor in Orthopaedic Surgery, University of Missouri, USA
Thomas Wild, Senior Consultant Surgeon, Wound Center, Clinic of Plastic, Hand and Aesthetic Surgery, University of Applied Sciences Anhalt; Clinic of Dermatology, Immunology and Allergology, Medical Center Dessau; Medical University Brandenburg, Theodor Fontane, Germany
DEFINING SURGICAL WOUND DEHISCENCE

The term ‘surgical wound dehiscence’ (SWD) can be interpreted by healthcare professionals in several ways. To some, SWD is reserved exclusively for the serious event of evisceration of abdominal contents that may occur following failure of a large abdominal surgical incision. But to others, the term has a broader meaning and covers a spectrum of problems ranging from superficial separation of part of an incision to complete separation of the full depth of the incision with exposure of body organs or surgical implants (Figure 1). This document considers SWD to apply to all degrees of separation of the margins of a closed surgical incision.

Research published on SWD has used a wide range of definitions. Variations in the definitions include:

- The term used for SWD (Box 1)
- Whether the definition relates to a surgical incision resulting from a specific type of surgery only (e.g. abdominal or cardiothoracic surgery) or to all types of surgery
- The tissue layers involved and/or the depth of the dehiscence
- The degree of dehiscence – i.e. involvement of part or the entire length of the incision
- The inclusion or exclusion of infected wounds
- Timing of the dehiscence in relation to surgery
- The need for a specific treatment – e.g. a further surgical procedure.

Some of the variation in definitions is due to the individual needs of the study and to aid extraction of data to answer the research question under investigation, e.g. data on a specific type of surgery or manifestation of SWD.

Box 1 | Synonyms for surgical wound dehiscence (SWD)

- Wound disruption
- Wound separation
- Wound opening
- Wound rupture
- Wound breakdown
- Wound failure
- Surgical site failure
- Post-operative wound dehiscence
- Burst abdomen
- Fascial dehiscence

Box 2 | Definition of SWD

Surgical wound dehiscence (SWD) is the separation of the margins of a closed surgical incision that has been made in skin, with or without exposure or protrusion of underlying tissue, organs or implants. Separation may occur at single or multiple regions, or involve the full length of the incision, and may affect some or all tissue layers. A dehisced incision may, or may not, display clinical signs and symptoms of infection.

N.B. Other types of closed wound may also dehisce, e.g. traumatic wounds that have been sutured. However, such wounds would not be considered to be SWD

There is currently no general standardised definition that aids understanding and accurate identification of SWD that can be used to underpin the principles of management. The Core Expert Working Group proposes a definition of SWD that can be applied to all closed surgical incision types (Box 2)
The causes of SWD can be categorised as:
- **Technical issues** with the closure of the incision – e.g. unravelling of suture knots
- **Mechanical stress** – e.g. coughing can cause breakage of the sutures or rupture of the healing incision after suture or clip removal/reabsorption
- **Disrupted healing** – e.g. due to comorbidities or treatments that hamper healing, or as a result of a surgical site infection (SSI) (Figure 2).

A wide variety of technical, mechanical and healing issues may contribute to SWD individually or in combination

**Technical issues**
SWD may occur because of technical issues with the closure of the incision. Surgical incisions are closed to bring together the sides of the wound to facilitate healing and minimise scar formation\(^\text{11,12}\). Surgical incision closure is achieved with sutures, staples, adhesive tapes or topical tissue adhesives. The most appropriate closure material and technique for a surgical incision depends on a wide variety of factors including the number of tissue layers to be closed, the anatomical location of the incision, the condition of the patient, and surgeon experience/preference\(^\text{13}\).

SWD may occur if the method of incisional closure fails or is not strong enough to hold the edges and sides of the incision together. For example, SWD may occur if suture knots slip or unravel, or sutures break, stretch, or cut through tissue because they have been placed too close to the edge of the incision, too far apart and/or put under too much tension\(^\text{14,15}\) (Figure 3). A retrospective study of 363 patients with SWD following laparotomy attributed 8% of SWD to broken sutures and 4% to loose knots\(^\text{16}\).

In addition to being caused by disrupted healing and mechanical stress, SWD can result from failure of the material used to close the incision, including stretching, slippage or breakage

**Mechanical stress**
Mechanical stress placed on a closed surgical incision can cause SWD by disrupting the material used for closure and/or rupturing the healing tissues (Box 3). Mechanical stress can result from excessive forced tension during wound closure or swelling of the tissues around the incision due to oedema. The latter may occur as part of the inflammatory phase of the healing process or in response to infection\(^\text{17}\) (Figure 4, page 7). Oedema may be an issue particularly for lower limb surgical incisions, e.g. following surgery for lower limb trauma\(^\text{18}\), and in patients with cardiac failure or who are critically ill and in fluid overload\(^\text{19}\).

Mechanical stress may also be due to a haematoma, seroma or abscess below the surface of the incision\(^\text{20}\).

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**Box 3 | Examples of causes of incisional mechanical stress that may result in SWD**
- Forced tension closure with inadequate tissue mobilisation or undermining
- Local oedema - e.g. due to inflammation, infection, position of the incision on a dependent anatomical area
- General oedema - e.g. in critical illness
- Incisional haematoma or seroma
- External trauma

**Abdominal or thoracic incisions**
- Increased intra-abdominal and/or intra-thoracic pressure - e.g. due to coughing, retching, vomiting, lifting heavy weights, abdominal compartment syndrome

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**Figure 2 | Causes of SWD** (adapted from\(^\text{10}\))

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**Figure 3 | SWD due to suture breakage**
In patients with abdominal and cardiothoracic incisions, mechanical stress may also arise from activities that cause a sudden increase in intra-abdominal and/or intra-thoracic pressure, e.g. retching, vomiting, coughing, sneezing and lifting heavy weights. Raised intra-abdominal pressure may also occur following abdominal surgery and, if sufficiently high, may compromise organ function (causing abdominal compartment syndrome) and contribute to SWD.

Patients should be advised to avoid placing undue stress on a closed surgical incision by following advice individually tailored according to patient factors and surgery type on: activity levels, avoiding overexertion, supporting/splinting the incision (e.g. with a surgical support bra or abdominal support), managing oedema and preventing trauma to the incision.

**Disrupted healing**

The complex process of wound healing in a closed surgical incision (known as healing by primary intention) can be divided into four distinct, necessary, but overlapping, phases: haemostasis, inflammation, proliferation and remodelling (Table 1).

Re-epithelialisation of a closed surgical incision is usually complete within 24–48 hours.

**Table 1 | Overview of the phases of wound healing of a surgical incision**

<table>
<thead>
<tr>
<th>Phase</th>
<th>Purpose</th>
<th>Timing after creation of surgical incision</th>
<th>Events</th>
</tr>
</thead>
<tbody>
<tr>
<td>Haemostatic</td>
<td>To prevent blood loss</td>
<td>Seconds to minutes</td>
<td>■ Cessation of bleeding through vasoconstriction, platelet aggregation and the release and activation of blood clotting factors to form a blood clot</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>■ Platelets release chemoattractants and growth factors for the recruitment of inflammatory cells</td>
</tr>
<tr>
<td>Inflammatory</td>
<td>To prevent infection and induce the proliferative phase of healing</td>
<td>Day 0 to up to several days</td>
<td>■ Vasodilatation and increased vascular permeability cause fluid leakage into the extravascular space (oedema/exudate.)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>■ Neutrophils are recruited to the wound site where they kill bacteria, degrade damaged or necrotic tissue and recruit other inflammatory cells such as macrophages</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>■ Macrophages and other immune cells support pathogen clearance and release a range of chemical factors that promote cell proliferation and synthesis of extracellular matrix</td>
</tr>
<tr>
<td>Proliferative</td>
<td>To repair the wound</td>
<td>Day 2 to up to several weeks</td>
<td>■ Fibroblasts migrate to the incision site and proliferate; collagen (especially type III) and extracellular matrix are synthesised; granulation tissue and new blood vessels are formed; keratinocytes migrate to re-epithelialise the wound</td>
</tr>
<tr>
<td>Remodelling/</td>
<td>To strengthen the repair</td>
<td>Day 21 to up to 2 years</td>
<td>■ Some type III collagen in the extracellular matrix is replaced by stronger type I collagen; myofibroblasts contract the wound to reduce scar surface area</td>
</tr>
<tr>
<td>maturation</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Table 2 | Tissue strength during healing**

<table>
<thead>
<tr>
<th>Time after incision</th>
<th>% of pre-incision breaking strength</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 week</td>
<td>3</td>
</tr>
<tr>
<td>3 weeks</td>
<td>30</td>
</tr>
<tr>
<td>3 months</td>
<td>80</td>
</tr>
</tbody>
</table>

**Table 3 | Proportion of dehisced wounds that are infected**

<table>
<thead>
<tr>
<th>Type of dehiscence</th>
<th>Proportion of dehisced wounds that are infected</th>
</tr>
</thead>
<tbody>
<tr>
<td>Abdominal dehiscence</td>
<td>52%–61%</td>
</tr>
<tr>
<td>Dehiscence following colorectal surgery</td>
<td>36.7%</td>
</tr>
<tr>
<td>Sternal incision dehiscence</td>
<td>49%</td>
</tr>
<tr>
<td>Episiotomy dehiscence</td>
<td>Up to 80%</td>
</tr>
</tbody>
</table>
SWD and other surgical site complications

Post-operative surgical site complications other than SWD include SSI, seroma, haematoma, delayed healing, poor quality or abnormal scar formation, and incisional hernia. Some surgical site complications increase the risk of SWD, e.g. SSI, seroma and haematoma. However, conversely, SWD increases the risk of SSI, delayed healing, poor quality scar formation and incisional hernia20,32 (Figure 5).

**SWD and SSI**

Infection occurs when microorganisms in a wound proliferate to a level that produces a local and/or systemic response33. Infection increases the production of degradative enzymes by immune cells and bacteria which can disrupt healing and weaken wound tissues16. As a result, SSI can cause SWD. This link between SWD and SSI is acknowledged in the Centers for Disease Control and Prevention definition of deep incisional SSI34 (Appendix 1, page 38). Conversely, however, not all infected incisions dehisce.

The link between SSI and SWD means that SSI can be a cause of and a risk factor for SWD

Although it is clear that some dehisced wounds are not due to infection (Table 3, page 6), rates of infection in dehisced wounds are infrequently reported35. In addition, infection can develop in a dehisced wound. Therefore, where infection rates are reported, it may not be clear whether infection occurred before or after dehiscence.

Unfortunately, some clinicians view SWD as synonymous with infection. In the age of awareness of the need for antimicrobial stewardship, accurate identification of infection in the context of SWD and the appropriate use of antimicrobials is ever more important.

Although there is a link between SWD and SSI, not all dehisced wounds are infected or require treatment for infection – and not all infected or inflamed wounds dehisce.

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**Box 4 | Examples of factors and conditions that may be associated with delayed or impaired wound healing27–31**

- **Local factors**
  - Hypoxia/ischaemia – e.g. due to peripheral arterial disease, oedema, respiratory disease
  - Devitalised tissue
  - Infection/contamination
  - Inflammatory conditions – e.g. pyoderma gangrenosum, vasculitis
  - Larger initial wound size
  - Ongoing mechanical stress or trauma
- **Systemic factors**
  - Advanced or very young age
  - Psychological stress
  - Chronic disease/comorbidities – e.g. diabetes mellitus, obesity, chronic kidney disease/uraemia, jaundice, chronic respiratory disease, immunosuppression
  - Medication – e.g. corticosteroids, chemotherapy
  - Radiotherapy
  - Smoking, alcoholism, substance misuse
  - Malnutrition
  - Connective tissue disorders – e.g. Ehlers-Danlos syndrome
  - Poor compliance with treatment plans
HOW COMMON IS SWD?

The difficulty of gaining a clear insight into the rates of occurrence of SWD is complicated by variations in the terminology used to described SWD (Box 1, page 4), the use of composite endpoints such as ‘wound complications’, and the lack of a generally accepted, standardised definition for SWD.

Under-reporting of SWD is also likely to occur for several other reasons including:
- Dehiscence, particularly of superficial, small areas of a wound, may not be recognised and recorded as SWD
- SWD may be overlooked and recorded as infection only, even when severe
- The trend for earlier discharge from hospital means that SWD is increasingly likely to occur in the community and may not be captured in hospital-based surveillance studies, and, particularly if relatively minor, may not be reported by patients or recognised by clinicians
- Negative implications for reimbursement and access to operating facilities may disincentivise reporting of surgical site complications.

Table 4 provides examples of SWD rates for different types of surgery.

There is considerable variation in SWD rates between surgical procedures, e.g. 0.65% for cardiothoracic surgery\(^38\) and 41.8% for pilonidal sinus surgery\(^39\)

A prospective study that analysed SWD rates following laparotomy by surgical wound class (i.e. clean, clean-contaminated, contaminated, or dirty or infected) reported that dehiscence was more common in the contaminated or dirty categories\(^40\) (Table 5).

In community settings, the most likely sites of SWD are the abdomen, leg and chest\(^53\)

### Table 4: Examples of SWD rates

<table>
<thead>
<tr>
<th>Surgical domain</th>
<th>Incidence</th>
</tr>
</thead>
<tbody>
<tr>
<td>Laparotomy(^9,36,44)</td>
<td>0.4%–3.8%</td>
</tr>
<tr>
<td>Cardiothoracic (sternotomy)(^3,38)</td>
<td>0.65%–2.1%</td>
</tr>
<tr>
<td>Orthopaedic surgery(^41–43)</td>
<td>1.1%–3.6%</td>
</tr>
<tr>
<td>Caesarean section(^14,54)</td>
<td>1.9%–7.6%</td>
</tr>
<tr>
<td>Oncoplastic breast reconstruction(^42,48)</td>
<td>4.6%–13.3%</td>
</tr>
<tr>
<td>Saphenous vein harvesting(^9)</td>
<td>8.9%</td>
</tr>
<tr>
<td>Pilonidal sinus (primary closure)(^39,50)</td>
<td>16.9–41.8%</td>
</tr>
<tr>
<td>Abdominoplasty following bariatric surgery(^51,52)</td>
<td>18.7%–21.5%</td>
</tr>
</tbody>
</table>

### Table 5: SWD rates following laparotomy according to surgical wound class\(^40\)

<table>
<thead>
<tr>
<th>Surgical wound category</th>
<th>Number of patients</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clean</td>
<td>0</td>
</tr>
<tr>
<td>Clean-contaminated</td>
<td>6 (12%)</td>
</tr>
<tr>
<td>Contaminated</td>
<td>22 (44%)</td>
</tr>
<tr>
<td>Dirty</td>
<td>22 (44%)</td>
</tr>
<tr>
<td>Total</td>
<td>50 (100%)</td>
</tr>
</tbody>
</table>
Impact of SWD

SWD can have a negative impact on patients’ mental health and physical and social functioning\(^5^4\)\(^5^5\). Mortality following sternal SWD can be very high (11%–53%), especially in the presence of infection or chronic obstructive pulmonary disease\(^3\)\(^5^6\). Mortality following abdominal SWD can also be high at 3%–35\%\(^5^4\)\(^5^7\). Furthermore, patients with abdominal SWD have a high risk of incisional hernia of up to 83\%\(^5^4\).

Analyses of data from US databases have illustrated the increased morbidity and mortality experienced by patients with SWD. An analysis of 2008 data from one database found that patients with SWD had in comparison with matched controls an additional:
- 9.6% mortality
- 9.4 days of hospitalisation
- US$40,323 of hospital charges\(^5^8\).

An analysis of 2003–2007 data from Veterans Health Administration reported that patients with SWD have a 61% higher odds of readmission within 30 days than patients without SWD\(^5^9\).

SWD can have a severe impact on a patient’s psychosocial wellbeing and carries considerable costs for healthcare systems

Costs and burden of SWD in the context of other wound types

It is becoming increasingly apparent that a considerable proportion of wounds with healing problems are surgical wounds and that these wounds are costly to manage. Contributors to costs may include frequent dressing changes, complications (such as infection) and hospital readmission\(^6^0\). Indirect costs may include loss of income, inability to participate in domestic or social activities, and welfare, social security or insurance payments\(^6^1\).

A recent US study of Medicare data from 2014 reported that:
- 14.5% of all Medicare beneficiaries were diagnosed with at least one type of wound or wound-related infection
- Infected surgical wounds were the most commonly treated wound type and affected 4% of all Medicare beneficiaries
- Costs for nonhealing and infected surgical wounds were the highest of any wound type (approximately US$13.1 billion), and greater than for the cost of treatment for diabetic foot ulcers
- There has been a considerable shift in total cost of care for all wounds from hospital inpatient to hospital outpatient settings, with outpatient costs about double inpatient costs\(^6^2\).

A UK study reported that more than half (57.1%) of wounds due to SWD healing by secondary intention, were being cared for in a community (rather than in a primary or secondary) setting\(^6^3\).

Furthermore, a study of the annual costs to the UK’s National Health Service (NHS) of caring for surgical wounds in a primary care setting reported that surgical wounds were the most costly and accounted for about 18.9%–21.8% of total expenditure on wound care\(^6^4\).

Data from large community-based organisations in Canada representing wound patients (n=24,678) have demonstrated that 43.9% of the wounds being managed are surgical wounds healing by secondary intention\(^6^5\). Patients with these wounds have required nursing care and clinical support over 6–69 weeks\(^6^5\).

A recent Australian study reported that the cost of management in a community nursing setting was AUD$509 for a patient with an uninfected SWD and AUD$1,025 with an infected SWD\(^5^3\).

SWD is not restricted to inpatient hospital care: it results in a high cost and resource burden in outpatient and community settings
RISK FACTORS FOR SWD

An understanding of the factors that increase a patient’s risk of SWD will guide the most appropriate prophylactic pre-, intra- and post-operative care.

The bulk of published research on risk factors specifically for SWD focuses on abdominal and sternal dehiscence with limited reporting across other surgical domains.

Table 6 lists general risk factors for SWD. The table differentiates between factors associated specifically with SWD and those that have often been reported as risk factors for SSI, haematoma or seroma (conditions that may themselves increase risk of SWD)

Major risk factors for SWD are obesity (body mass index (BMI) ≥35kg/m²), diabetes mellitus, current or recent smoking, emergency surgery, age >65 years, extended duration of surgery, inadequate surgical closure, peri-operative hypothermia and wound infection.

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**Table 6 | Main general risk factors for SWD**

<table>
<thead>
<tr>
<th>Category of risk factor</th>
<th>Patient-related modifiable risk factors</th>
<th>Pre-operative risk factors</th>
<th>Intra-operative risk factors</th>
<th>Post-operative risk factors</th>
</tr>
</thead>
<tbody>
<tr>
<td>Major</td>
<td>BMI ≥35.0kg/m²</td>
<td>Emergency surgery</td>
<td>Extended duration of surgery</td>
<td>Wound infection (SSI)</td>
</tr>
<tr>
<td></td>
<td>Diabetes mellitus</td>
<td>Age &gt;65 years</td>
<td>Inadequate surgical closure</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Current or recent smoking</td>
<td></td>
<td>Perioperative hypothermia*</td>
<td></td>
</tr>
<tr>
<td>Moderate</td>
<td>COPD‡</td>
<td>Male gender</td>
<td>Blood transfusion</td>
<td>Failure to wean from ventilator</td>
</tr>
<tr>
<td></td>
<td>Malnutrition: hypoalbuminaemia</td>
<td>ASA Physical Status ≥2</td>
<td>Junior surgeon</td>
<td>One or more complication other than dehiscence</td>
</tr>
<tr>
<td></td>
<td>(serum albumin &lt;3.0g/dL)</td>
<td>Previous dehiscence/</td>
<td>High wound tension closure</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Anaemia</td>
<td>wound healing problems</td>
<td>Tissue trauma/ large area of dissection and/ or undermining</td>
<td></td>
</tr>
<tr>
<td></td>
<td>BMI 30.0–35.0kg/m²</td>
<td>Immunosuppression</td>
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<td></td>
<td>Alcohol abuse</td>
<td>Long-term steroid use</td>
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<td></td>
<td></td>
<td>Malignant disease</td>
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<td></td>
<td></td>
<td>Chemotherapy</td>
<td></td>
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<td></td>
<td></td>
<td>Radiotherapy</td>
<td></td>
<td></td>
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<tr>
<td></td>
<td></td>
<td>Uraemia</td>
<td></td>
<td></td>
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<tr>
<td></td>
<td></td>
<td>Peripheral vascular disease</td>
<td></td>
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<tr>
<td></td>
<td></td>
<td>Suboptimal timing or omission of prophylactic antibiotics*</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Minor</td>
<td>BMI 25.0–29.9kg/m²</td>
<td>Extended pre-operative hospitalisation or residency in a nursing home*</td>
<td>Failure to obliterate dead space</td>
<td>Trauma across incision</td>
</tr>
<tr>
<td></td>
<td>Congestive cardiac failure</td>
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<tr>
<td></td>
<td>Cardiovascular disease</td>
<td></td>
<td></td>
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<tr>
<td>Rare</td>
<td>Alpha-1 antitrypsin deficiency</td>
<td></td>
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<tr>
<td></td>
<td>Ehler-Danlos syndrome</td>
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<tr>
<td></td>
<td>Behçet’s disease</td>
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<tr>
<td></td>
<td>Bleeding disorders*</td>
<td></td>
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<td></td>
</tr>
</tbody>
</table>

‡ May be a risk factor in different types of surgery for different reasons, e.g. because of coughing in abdominal surgery and sternotomy and because of the adverse effects of chronic disease on wound healing in all types of surgery.

* These are risk factors for SSI or other surgical wound complications, e.g. haematoma and seroma, that may be associated with SWD. Other factors listed in the table have been reported to be associated with SWD specifically.

ASA: American Society of Anesthesiologists; COPD: chronic obstructive pulmonary disease; SSI: surgical site infection.
<table>
<thead>
<tr>
<th>Type of surgery</th>
<th>Additional risk factors</th>
</tr>
</thead>
<tbody>
<tr>
<td>Abdominal</td>
<td>See Table 6, page 10, for general risk factors for SWD</td>
</tr>
<tr>
<td></td>
<td>■ Midline laparotomy</td>
</tr>
<tr>
<td></td>
<td>■ Damage to the gastrointestinal tract</td>
</tr>
<tr>
<td></td>
<td>■ Intestinal or biliary tract surgery</td>
</tr>
<tr>
<td></td>
<td>■ Creation of an ostomy</td>
</tr>
<tr>
<td></td>
<td>■ Muscle flap creation</td>
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<tr>
<td></td>
<td>■ Loss of visceral domain &gt;20%</td>
</tr>
<tr>
<td></td>
<td>■ Peritonitis</td>
</tr>
<tr>
<td></td>
<td>■ Sepsis</td>
</tr>
<tr>
<td></td>
<td>■ Jaundice</td>
</tr>
<tr>
<td></td>
<td>■ Ascites</td>
</tr>
<tr>
<td></td>
<td>■ Coughing/pulmonary problems/pneumonia</td>
</tr>
<tr>
<td></td>
<td>■ Post-operative anastomotic dehiscence/fistula</td>
</tr>
<tr>
<td></td>
<td>■ CVA without residual deficit</td>
</tr>
<tr>
<td>Breast/plastic</td>
<td>See Table 6, page 10, for general risk factors for SWD</td>
</tr>
<tr>
<td></td>
<td>■ SWD element of the Breast Reconstruction Risk Assessment (BRA) score</td>
</tr>
<tr>
<td></td>
<td>■ Previous surgery at same site</td>
</tr>
<tr>
<td>Cardiothoracic</td>
<td>See Table 6, page 10, for general risk factors for SWD</td>
</tr>
<tr>
<td></td>
<td>■ Osteoporosis</td>
</tr>
<tr>
<td></td>
<td>■ Antiplatelet medication</td>
</tr>
<tr>
<td></td>
<td>■ Previous sternotomy</td>
</tr>
<tr>
<td></td>
<td>■ Prolonged cardiopulmonary bypass time</td>
</tr>
<tr>
<td></td>
<td>■ Chronic cough</td>
</tr>
<tr>
<td></td>
<td>■ NYHA functional class IV</td>
</tr>
<tr>
<td></td>
<td>■ Bilateral internal mammary artery harvest</td>
</tr>
<tr>
<td></td>
<td>■ Post-operative pneumonia</td>
</tr>
<tr>
<td></td>
<td>■ Beta-blocker use</td>
</tr>
<tr>
<td></td>
<td>■ Previous surgery in current admission</td>
</tr>
<tr>
<td></td>
<td>■ Respiratory failure</td>
</tr>
<tr>
<td></td>
<td>■ Urinary tract infection</td>
</tr>
<tr>
<td></td>
<td>■ Left ventricular assist device*</td>
</tr>
<tr>
<td></td>
<td>■ Transplant*</td>
</tr>
<tr>
<td></td>
<td>■ Cardiopulmonary bypass time extended*</td>
</tr>
<tr>
<td>Obstetric</td>
<td>See Table 6, page 10, for general risk factors for SWD</td>
</tr>
<tr>
<td></td>
<td>■ Human papilloma virus (HPV) infection</td>
</tr>
<tr>
<td>Episiotomy repair:</td>
<td>■ African-American race</td>
</tr>
<tr>
<td>Caesarean section:</td>
<td>■ Vertical incision</td>
</tr>
<tr>
<td></td>
<td>■ Stapled wound closure</td>
</tr>
<tr>
<td></td>
<td>■ Chorioamnionitis</td>
</tr>
<tr>
<td></td>
<td>■ Multiple caesarean sections*</td>
</tr>
<tr>
<td></td>
<td>■ Operative blood loss &gt;1.5l*</td>
</tr>
<tr>
<td></td>
<td>■ Pre-eclampsia</td>
</tr>
<tr>
<td>Orthopaedic</td>
<td>See Table 6, page 10, for general risk factors for SWD</td>
</tr>
<tr>
<td></td>
<td>■ Implant-related surgery</td>
</tr>
<tr>
<td></td>
<td>■ Poor compliance by patients with nurses’ recommendations</td>
</tr>
<tr>
<td></td>
<td>■ Traumatic injury</td>
</tr>
<tr>
<td></td>
<td>■ Rheumatoid arthritis*</td>
</tr>
</tbody>
</table>

* Risk factors for SSI or other surgical wound complications, e.g. haematoma and seroma, that may be associated with SWD. Other factors listed in the table have been reported to be associated with SWD specifically.

ASA: American Society of Anesthesiologists; CVA: cerebrovascular accident; NYHA: New York Heart Association functional class.
IDENTIFYING SWD

SWD dehiscence can occur at any time after surgery, from one day to more than 20 days after surgery, but generally occurs at post-operative days 4–14.

Monitoring the healing progress of a surgical incision will enable the identification of incisions in which healing is progressing well and those in which healing is impaired and has the potential to progress to SWD (Table 8).

Signs of probable SWD

SWD can occur without warning. Incisions at risk of dehiscence may show signs of inflammation beyond the time and extent expected for normal healing, e.g. more exaggerated incisional redness, swelling, warmth and pain that extend beyond post-operative day 5. Palpation of the incision and surrounding area may reveal warmth and a collection of fluid under some or all of the incision (a seroma, haematoma or abscess). A sudden increase in pain or discharge of serosanguineous fluid from the incision may herald SWD.

Signs of inflammation at the incisional site, e.g. warmth, erythema, oedema, discolouration and pain, are normal during the first few days after surgery, and do not necessarily indicate infection.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Relationship to TIME framework*</th>
<th>Signs that incisional healing is progressing well</th>
<th>Signs that healing is impaired</th>
</tr>
</thead>
</table>
| Incision colour | Tissue | - Days 1–4: red  
- Days 5–14: bright pink  
- Day 15–1 year: pale pink, progressing to white or silver in light-skinned patients or to darker than usual skin colour in patients with darkly-pigmented skin | - Days 1–4: may be red, tension in the incision line  
- Days 5–9: edges may be well-approximated and the tension remains  
- Days 10–14: if SWD does not occur, colour may remain red or progress to pink and may be followed ultimately by hypertrophic scarring |
| Healing ridge | - Days 5–9: a healing ridge of thickened tissue indicating newly formed collagen can be felt about 1cm either side of the incision along its length, and persists into the remodelling phase | Lack of healing ridge |
| Peri-incisional area | Infection/inflammation | - Signs of inflammation:  
- Mild oedema, erythema, warmth or skin discolouration that resolves by day 5  
- Pain | - Signs of inflammation may be absent in the first few days after surgery  
- Signs of inflammation and ongoing pain may be present for extended periods |
| Exudate | Moisture | - Days 1–4: decreasing in volume from moderate to minimal and changing from sanguineous (blood) to serosanguineous (mixture of blood and serum) to serous (clear, amber serum)  
- Resolves by day 5 | - Exudate persists beyond days 1–4  
- Exudate may be serosanguineous, serous or purulent (e.g. cloudy, green, yellow or brown) |
| Wound margins | Edge | - Epithelial closure should be seen by day 4 along the entire incision  
- Approximated | - Epithelial resurfacing may be only partially present or entirely absent  
- Area(s) of separation (SWD) may be present by day 14 |

*See Table 9, page 16, for more information about using the TIME framework in the assessment of SWD
Signs of SWD
Areas of separation of the wound margins may vary from tiny 'pin pricks' to larger gaping areas to the entire length and depth of the incision. If the incision opens into a body cavity, SWD may result in evisceration. Sutures or clips may be visible in the separated area and may be broken.

In patients with abdominal or sternal incisions, dehiscence may follow an episode of retching, vomiting or coughing. Patients may describe a sensation of pulling or ripping in the area of the incision, or the feeling that something has given way.

Signs of infection
A patient with a surgical incision at increased risk of SWD or that has dehisced can show local and systemic signs and symptoms of infection (Box 5 and Box 6).

The diagnosis of infection (SSI) in surgical incisions or SWD should be made on the basis of clinical signs and symptoms

Box 5 | Local clinical signs and symptoms of infection in a closed surgical incision

See Box 6 for systemic signs and symptoms of infection
- Erythema - localised or spreading (cellulitis)
- Pus/purulent or haemopurulent exudate
- Abscess
- Swelling/induration
- Local warmth
- Malodour
- Crepitus (crackling feeling or sound detected on palpation due to gas in the soft tissues)
- Dehiscence
- Unexpected pain or tenderness

Box 6 | Systemic signs and symptoms that may be associated with infection of a closed or dehisced surgical incision

- Malaise
- Loss of appetite
- Pyrexia or hypothermia
- Tachycardia
- Tachypnoea
- Elevated C-reactive protein (CRP)
- Elevated or suppressed white blood cell count
- Sepsis
- Septic shock

Various systems exist to aid diagnosis of SSI. These include the CDC definitions of SSI and adaptations such as the definitions used by Public Health England (PHE) (Appendix 1, page 38), and the ASEPSIS scoring system. The ASEPSIS system is an objective means of assessing surgical incisions for infection and results in a numerical score that indicates the presence and severity of any infection (Appendix 2, page 39).
Patients with SWD of any depth or length should receive a structured holistic assessment that includes assessment of the general condition of the patient and of the dehisced incision.

Assessment of a patient with SWD will provide important information that will guide management (Figure 6, page 15), including:
- Modifiable factors that may be hindering healing
- Any signs of local or systemic infection
- Whether further investigations are required
- The condition of the dehisced area.

The results of the holistic assessment, which should be fully documented, will guide the most appropriate management.

General assessment
Box 7 outlines the components of a general assessment of a patient with SWD, which includes all facets of previous and current health and psychosocial status.

The main aims of general assessment are to identify any factors that may have contributed to or exacerbate the dehiscence or that may impair healing, and to detect any clinical signs of systemic infection.

Box 7 | General assessment of a patient with SWD (adapted from 96)
See Tables 6 and 7 (pages 10 and 11) for details of risk factors for SWD

- Medical and surgical history, including:
  - Previous problems with wound healing – e.g. SWD, SSI
  - Radiotherapy
  - Chemotherapy
  - Allergies and sensitivities to medication and skin/wound products

- Nature of the surgical procedure that resulted in the incision that has dehisced, including:
  - Reason for surgery and date
  - Emergency/elective
  - Intra-operative and post-operative complications – e.g. haemorrhage, hypothermia, duration of surgery, SSI
  - Closure method
  - Date of suture/clip removal

- Current health, including:
  - Need for haemodynamic or ventilatory support
  - Active comorbidity – e.g. diabetes mellitus, obesity, COPD, blood clotting factor deficiencies, anaemia/blood transfusions, cough/chest infection; constipation, dermato logical conditions
  - Nutritional status – e.g. presence of malnutrition, level of hydration, ability to eat and drink
  - Physical parameters relating to possible systemic infection – e.g. core temperature, levels of inflammatory markers (e.g. CRP) and white blood cell (WBC) count

- Lifestyle, including smoking, alcohol intake, diet, level of physical activity

- Current medication and reasons for use, including:
  - Anticoagulant/antiplatelet treatment
  - Chronic corticosteroids
  - Immunosuppressants
  - Antibiotics
  - Analgesics

- Pain, including current location and severity of pain, whether related to the wound or elsewhere; use of numeric or visual analogue scales can aid objective assessment and monitoring of pain severity; current pain management

- Psychosocial status, including:
  - Care setting
  - Family/carer support
  - Occupation and financial situation
  - Patient’s understanding of and attitude to their condition and the incision and surgery
  - Ability and willingness to engage in care
  - Impact of wound on quality of life (physical, social and emotional)

*To calculate number of days since surgery; very early dehiscence may be due to technical issues and duration of SWD may influence management
†Of particular relevance in patients which cardiothoracic or abdominal incisions
‡Post-operative mobilisation is important, however, depending on the position of the wound, overexertion may contribute to or exacerbate SWD
Incision/wound assessment
Prior to assessment of SWD, the events, if any, leading to the dehiscence, e.g. coughing, vomiting, trauma, suture/clip removal, purulent drainage, should be ascertained. The duration of the dehiscence should also be determined: SWD occurring very soon after surgery and of very recent occurrence may be suitable for re-suturing.

The entire length of an incision with SWD should be fully assessed: the factors that led to the SWD may also be affecting other regions of the incision that remain closed.

The Core Expert Working Group recommends the use of a structured framework, e.g. TIME, to aid assessment of SWD (Table 9, page 16). Sequential photographs can aid monitoring. Photographs should be obtained and stored after gaining patient consent and according to local policy.

If more than one area of dehiscence is present, each area should be assessed individually (Figure 7)

A short area of dehiscence is not necessarily only superficial and may extend deeply

While it is important to determine the depth of an area of dehiscence, any probing should be undertaken very gently and carefully by a clinician with suitable competency to avoid inadvertently exacerbating the dehiscence or causing other damage

All general and wound assessments, further tests, interventions and referrals should be documented
Table 9 | Assessment of SWD using the TIME framework (adapted from93,96,97,99)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Assess Specifics</th>
</tr>
</thead>
</table>
| **Tissue**               | Location and extent of dehiscence  
  ■ Location of the incision  
  ■ Proportion of the incision affected  
  ■ Number of areas of dehiscence  
  ■ Presence of sutures/clips and condition (intact/broken)  
  Depth of dehiscence  
  ■ Partial or full-thickness dehiscence and tissue layers affected (see Figure 8, page 18); WUWHS SWD Grade (see Table 10, page 18)  
  ■ Extension to or exposure of organs/bone/implant  
  ■ Presence of undermining/tunnelling  
  ■ For abdominal SWD, presence of evisceration  
  Tissue viability  
  ■ Condition of exposed tissues  
  ■ Wound bed tissue types and proportions – e.g. of necrotic/devitalised tissue, slough and granulation tissue  
  Dimensions  
  ■ Dimensions of the dehisced area(s): maximum length, width, depth  
| **Infection (or inflammation)** | For local indicators of infection or inflammation  
  ■ Clinical signs and symptoms  
  ■ See Box 5 and Box 6, page 13, and Box 8, page 17, for signs and symptoms of acute and chronic infection  
  ■ N.B. In patients who are immunosuppressed, signs and symptoms may be less obvious  
  Moisture | Exudate/drainage colour, consistency, type and odour  
  ■ Purulent (cream, yellow or green) or haemopurulent (red, brown) may indicate infection  
  ■ Yellow or brown exudate may indicate a urinary or enteric fistula  
  ■ Malodour may indicate infection or fistula  
  Exudate/drainage level  
  ■ Indications of the level of exudate production can be gained from the condition of the current dressing (i.e. a dry dressing indicates low exudate levels; a saturated or leaking dressing indicates higher levels) and the appearance of the wound bed  
| **Edge**                 | Edges of dehisced area  
  ■ In long-standing areas of dehiscence, the edges may become undermined  
  Colour and condition of the surrounding skin  
  ■ Signs of dermatological conditions that may affect healing – e.g. radiation dermatitis  
  ■ Signs of spreading infection – e.g. spreading erythema, warmth and oedema  
  ■ Periwound maceration may indicate high exudate/drainage levels and/or inadequate absorbency of the dressing  

**Diagnosis of infection**

The diagnosis of infection of a surgical incision or a dehisced wound is largely based on local and systemic clinical signs and symptoms (Box 5 and Box 6, page 13, and Box 8, page 17). Fever in the first 48 hours after surgery is unlikely to be due to SSI. The role of sampling and microbiological culture in the diagnosis of SSI continues to be debated. Reasons for this include that superficial sampling, such as swabbing, may reflect only surface bacteria and not bacteria in deeper tissues – an issue of particular relevance to deep surgical wounds. Technological developments, such as the use of point-of-care fluorescence imaging (e.g. MolecuLight iX™, distributed by Smith & Nephew) to detect areas of tissue with increased bacterial levels and guide sampling, may help to increase the usefulness of microbiological sampling.

Clinicians should be aware of the limitations of microbiological analysis of wound samples, and should interpret the results in the context of clinical signs and symptoms, noting that a ‘negative’ swab does not necessarily exclude infection.

**Imaging diagnostics**

Most patients with SWD do not require further investigation with imaging diagnostics. However, if there is uncertainty about the diagnosis, the depth of dehiscence, or if an area of dehiscence is increasing in size or is not improving despite treatment, imaging may be warranted.

In many cases, ultrasound scanning will be the most appropriate imaging modality, with more expensive modalities such as magnetic resonance imaging (MRI) reserved for further
investigation. In addition to assessing the tissues, imaging may be used to detect and assess seromas, haematomas and collections of pus, and to evaluate the proximity of the dehiscence to implants such as meshes or prosthetic joints.

Grading of SWD

Systems for grading or classifying SWD often relate to specific types of surgery, e.g. thoracic or abdominal surgery. Some classifications are adaptations of the adverse event reporting systems, e.g. of the Ottawa Thoracic Morbidity and Mortality system or the Clavien-Dindo system.

There is a need for a general classification system for SWD that is applicable to incisions from all surgery types, is easy to use, is suitable for use in all care settings (including community settings), that indicates severity, and that can be linked to appropriate management strategies.

The proposed WUWHS SWD Grading System in Table 10, page 18, was developed by the Core Expert Working Group during the consensus meeting and is an adaptation of the Sandy SWD Grading system.

The system uses depth and the presence of infection as the main determinants of SWD severity. Distinguishing SWD with no clinical signs and symptoms of infection from SWD with clinical signs and symptoms of infection is intended to emphasise the differences in approach to management that may be required.

Assignment of a WUWHS SWD Grade should take place only after full assessment of the patient and the surgical incision, including probing and exploration of the areas of dehiscence if required by a clinician with suitable competency.

Even though most SWD occurs 4–14 days post-operatively, a time-period of 30 days has been included in the grading system. The inclusion of a time-period is intended to encourage surveillance and reporting of SWD post-discharge as, in common with SSI, the probable under-reporting of SWD may be related to occurrence of SWD after a patient has left hospital. The time-period of 30 days is broadly in line with reporting requirements for SSI and has been applied to all SWD grades for consistency. As more is learnt about SWD, the time-period may need to be adjusted.

As SWD generally occurs at days 4-14 post-operatively, a significant proportion is likely to occur after discharge.

Figure 9, page 19, illustrates how the tissue layers relate to the WUWHS SWD grading in Table 10, page 18.

No matter how long the area of dehiscence is, SWD involving the deep layers of an incision is more serious than that involving more superficial layers.

---

**Box 8 | Clinical signs and symptoms of local wound infection in a chronic SWD**

See Box 6, page 13, for systemic signs and symptoms of infection

- New, increased, or altered pain*
- Delayed healing*
- Malodour or change in odour
- Increased or altered/purulent exudate
- Periwound oedema
- Bleeding or easily damaged granulation tissue
- Altered wound bed colour
- Induration of periwound skin
- Pocketing and bridging

*Individually highly indicative of infection. Infection is highly likely in the presence of two or more of the signs above
### Table 10 | Proposed WUWHS SWD Grading System (adapted from Sandy SWD Grading System\(^{100}\))

**Definition:** Surgical wound dehiscence (SWD) is the separation of the margins of a closed surgical incision that has been made in skin, with or without exposure or protrusion of underlying tissue, organs or implants. Separation may occur at single or multiple regions, or involve the full length of the incision, and may affect some or all tissue layers. A dehisced incision may, or may not, display clinical signs and symptoms of infection.

<table>
<thead>
<tr>
<th>WUWHS SWD Grade*</th>
<th>Descriptors</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>1</strong> Figure 9a, page 19</td>
<td>Dermal layer only involved; no visible subcutaneous fat</td>
</tr>
<tr>
<td>1a Figure 9b, page 19</td>
<td>As Grade 1 plus clinical signs and symptoms of infection (e.g. superficial incisional SSI(^{\S}))</td>
</tr>
<tr>
<td><strong>2</strong> Figure 9c, page 19</td>
<td>Subcutaneous layer exposed; fascia not visible</td>
</tr>
<tr>
<td>2a Figure 9d, page 19</td>
<td>As Grade 2 plus clinical signs and symptoms infection (e.g. superficial incisional SSI(^{\S}))</td>
</tr>
<tr>
<td><strong>3</strong> Figure 9e, page 19</td>
<td>Subcutaneous layers and fascia exposed</td>
</tr>
<tr>
<td>3a Figure 9f, page 19</td>
<td>As Grade 3 plus clinical signs and symptoms infection (e.g. deep incisional SSI(^{\S}))</td>
</tr>
<tr>
<td><strong>4</strong> Figure 9g, page 19</td>
<td>Any area of fascial dehiscence with organ space, viscera, implant or bone exposed</td>
</tr>
<tr>
<td>4a Figure 9h, page 19</td>
<td>As Grade 4 plus clinical signs and symptoms infection (e.g. organ/space SSI(^{\S}))</td>
</tr>
</tbody>
</table>

*Grading should take place after full assessment including probing or exploration of the affected area as appropriate by a clinician with suitable competency
†Where this is >1 region of separation of the wound margins, SWD should be graded according to the deepest point of separation
‡Where day 1 = the day of the procedure
§See Appendix 1, page 38, for the CDC definitions of the different types of SSI
^Grade 4/4a dehiscence of an abdominal incision may be called ‘burst abdomen’

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**Figure 8 | Proposed WUWHS SWD Grade according to the tissue layers involved in the dehiscence**

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**Table 10 | Proposed WUWHS SWD Grading System (adapted from Sandy SWD Grading System\(^{100}\))**

<table>
<thead>
<tr>
<th>Tissue layers</th>
<th>WUWHS SWD Grade</th>
</tr>
</thead>
<tbody>
<tr>
<td>Skin</td>
<td>Grade 1</td>
</tr>
<tr>
<td>Subcutaneous tissues</td>
<td>Grade 2</td>
</tr>
<tr>
<td>Muscle</td>
<td>Grade 3</td>
</tr>
<tr>
<td>Deep fascia</td>
<td>Grade 4</td>
</tr>
<tr>
<td>Organ/implant</td>
<td></td>
</tr>
</tbody>
</table>
Figure 9 | Examples of the proposed WUWHS SWD Grades

a) WUWHS SWD Grade 1
Small area of dermal separation

b) WUWHS SWD Grade 1a
Post-mastectomy: small areas of dermal separation with inflammation and infection

c) WUWHS SWD Grade 2
Obese patient with exposed subcutaneous tissue and tunnel into pannus following surgery for seatbelt trauma

d) WUWHS SWD Grade 2a
Post-mammaplasty: dermal separation with exposure of subcutaneous tissue with inflammation and purulent exudate

e) WUWHS SWD Grade 3
Post-spinal surgery: full length dehiscence with fascial exposure without signs of infection

f) WUWHS SWD Grade 3a
Leg incision: dehiscence exposing muscle and fascia with pus and cellulitis

g) WUWHS SWD Grade 4
Post-laparotomy: dehiscence with abdominal organ exposure and no signs of infection

h) WUWHS SWD Grade 4a
Separation of suture line with exposed hardware with inflammation and signs of infection
MANAGEMENT OF SWD

SWD can vary from a shallow area of a small proportion of an incision to the full depth of an entire incision with eversionation of organs or exposure of an implant. Even so, the goal of SWD management is usually closure of the wound.

The management of SWD should be tailored to the individual patient and often requires involving and working collaboratively with the patient, family, carers and wider multidisciplinary team.

Before planning management, it is essential that the clinician has a clear understanding of the structures (e.g. implants, vital organs or bone) located directly beneath the dehisced wound to ensure correct management and to avoid exacerbating the patient’s condition or causing a more serious complication.

The principles of management of SWD include:

- Reassurance, management of expectations and education
- Pain management
- Removal or amelioration of risk factors that may have contributed to SWD or that may compromise healing
- Management of systemic infection
- Local management of the dehisced wound, including management of local infection.

The objectives of treatment and the management plan should be fully documented and discussed with the patient, carers and family.

Reassurance, management of expectations and patient education

SWD is potentially frightening for patients, even if a relatively small proportion of the incision is involved. Patients will need to be reassured with an explanation tailored to their needs and understanding of what has happened, the possible reasons for it happening, the actions to be taken and the longer-term outlook. Patients should be encouraged to voice any concerns and may find it valuable to talk to a patient who has experienced similar issues.

Education of a patient with SWD should include signs and symptoms of infection (if not already present), how to avoid putting additional stress on the incision, advice about activity levels, and individualised instructions on what to do and who to contact if the condition of the wound or patient deteriorates.

Pain management

Pain management should include management of background pain and pain associated with dressing/device changes and debridement. Pharmacological and non-pharmacological measures should be considered, including education and careful selection of dressings, change frequency and change technique to minimise pain and trauma. The World Health Organization’s three-step cancer pain ladder for adults can be applied to the management of pain in other contexts and may be useful in guiding appropriate pharmacological therapy (www.who.int/cancer/palliative/painladder/en/).

Management of comorbidities and contributory factors

Any modifiable factors that might have contributed to SWD or that may impede healing, e.g. chest infection, poor blood glucose control in patients with diabetes mellitus, smoking and inadequate nutrition, should be addressed.
Local management of SWD

The local management of SWD is dependent on a range of factors including the:

- **Severity of the dehiscence** – e.g. depth/WUWHS SWD Grade and exposure of viscera, bone or implants
- **Presence of infection**
- **Timing of the dehiscence** in relation to the surgery that produced the incision
- **Presence of comorbidities** that increase the risk of surgical site complications and/or impair healing.

The results of the holistic assessment will indicate appropriate treatment objectives and guide management planning.

Local management of a dehisced wound can be guided by application of the TIME Framework (Table 9, page 16) with consideration of removal of non-viable tissue (debridement), management of infection, exudate control and promotion of moist wound healing.
Figure 11 | Local management of a dehisced wound according to WUWHS SWD Grade

See Table 10, page 18, for the WUWHS SWD Grading System, and the vignettes on page 31 for examples of management of different grades of SWD.

<table>
<thead>
<tr>
<th>Assessment</th>
<th>Management objectives</th>
<th>General management</th>
<th>Local management</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>See Figure 10, page 21</td>
<td></td>
</tr>
</tbody>
</table>

### GRADE 1

**Usually secondary***

- If needed, often autolytic
- Apply dressing appropriate for exudate level, periwound skin condition and need for autolytic debridement
- Consider NPWT if patient is at risk of delayed healing***

### GRADE 2

**Usually secondary***

- Often autolytic; +/- other methods if there is extensive non-viable tissue
- Apply dressing(s) +/- wound fillers appropriate for depth of wound, exudate level, periwound skin condition and need for autolytic debridement
- Consider NPWT if patient has high exudate levels or is at risk of delayed healing***, choosing device and wound fillers/wound contact layers according to exudate level and depth of wound

### GRADE 3

**Often secondary**: delayed primary if contamination or healing problems; +/- flaps, skin grafts, dermal matrices/substitutes**

- Sharp/surgical debridement to remove necrotic material
- Consider use of a prophylactic antimicrobial dressing in patients at high risk of delayed healing***
- Apply dressing(s) +/- wound fillers appropriate for depth of wound, exudate level and periwound skin condition
- Consider NPWT if patient has high exudate levels or is at risk of delayed healing***, choosing device and wound fillers/wound contact layers according to exudate level and depth of wound

### GRADE 4

**Usually delayed primary*** +/− sequential layered closure, flaps, skin grafts, dermal matrices/substitutes**

- Surgical debridement to remove necrotic/non-viable tissue +/- lavage
- +/- Surgical management of viscera, bone or implants as appropriate
- Consider NPWT with wound fillers +/- prophylactic antimicrobial interface +/- instillation/irrigation

### Clinical signs or symptoms of infection

#### GRADE 1a

- As for 1 above and:
  - Manage local infection: apply antimicrobial dressing according to best practice guidelines and local formulary
  - Manage systemic infection, if present§

#### GRADE 2a

- As for 2 above and:
  - Manage local infection: apply antimicrobial dressing according to best practice guidelines and local formulary
  - If using NPWT, consider an antimicrobial interface according to local protocol
  - Manage systemic infection, if present§

#### GRADE 3a

- As for 3 above and:
  - If using NPWT, consider an antimicrobial interface according to local protocol
  - Manage systemic infection, if present§

#### GRADE 4a

- As for 4 above and:
  - +/- Surgical management of viscera, bone or implants if a source of ongoing infection
  - If using NPWT, consider an antimicrobial interface according to local protocol +/− instillation/irrigation
  - Manage systemic infection, if present§

### Monitoring and reassessment

See Figure 10, page 21

*Primary closure may be appropriate if SWD occurs <48 hours after surgery for technical reasons and is not otherwise contraindicated

**Dermal matrices/substitutes should not be used in the presence of wound infection

***See Box 4, page 7, for examples of factors that may delay healing

§Manage systemic infection according to best practice guidelines, taking into account local policies and results of any microbiological culture and sensitivity reports
Method of closure
An important initial decision in the management of SWD is about the most appropriate method for achieving closure of the wound. This will largely depend on timing in relation to the surgery that produced the incision, the depth of the dehiscence (i.e. WUWHS SWD Grade), the location of the incision and whether infection is present.

Primary closure
Primary closure following SWD (Figure 12) may be indicated if:
- SWD has occurred within 48 hours of surgery and is clearly the result of a technical issue, e.g. sutures have come undone, clips were not properly applied
- No other problems have contributed to the SWD – i.e. there is no undue tension on the incision and there are no signs of infection
- The patient is not at increased risk of surgical site complications.

Secondary closure
Secondary closure (Figure 12) is frequently used in superficial SWD with or without infection, e.g. WUWHS SWD Grades 1, 1a, 2 and 2a. It may also be used in deeper dehiscence, e.g. WUWHS SWD Grades 3 and 3a, and occasionally WUWHS SWD Grades 4 and 4a, where there is a high risk of SSI, infection is present, or where primary closure is not possible, e.g. because of tissue loss.

Delayed primary closure
Delayed primary closure (Figure 12), sometimes referred to as healing by tertiary intention, is mainly used in the management of deeper SWD, e.g. WUWHS SWD Grades 3, 3a, 4 and 4a, where the incision is contaminated or infected, or where the risk of recurrence of dehiscence is high because of comorbidities or subcutaneous/visceral swelling that would put tension on a resutured incision. When the time for primary closure arrives, a flap or skin graft may be used if a tissue defect remains.
Management of abscess, seroma and haematoma
The collection of fluid, whether pus, serous fluid or blood, under a closed incision may increase incisinal tension and the risk of SWD. Abscesses (Figure 13) should be drained to remove pus and a potential source of ongoing infection. Seromas and haematomas may resolve spontaneously. However, depending on the size, location and impact on the incision, seromas and haematomas may require aspiration or the insertion of a drain.

Debridement
Necrotic and non-viable tissue and foreign material in a dehisced incision can act as culture media and foci for bacterial growth and the formation of biofilm and so increase the risk of infection and impaired healing. The presence of microbial biofilm in the incision may be related to up to 80% of SSIs. Debridement removes non-viable tissue and foreign material, reducing bioburden, biofilm and inflammatory stimulus. Particularly in sharp or surgical debridement, debridement also stimulates the release of growth factors involved in healing.

There are several methods of debridement (Table 11). Clinicians should work within the limits of their competency when conducting debridement and refer the patient on if a debridement method beyond their competency is required.

Autolytic debridement is often sufficient for dehisced incisions graded as WUWHS SWD Grades 1/1a and 2/2a. Sharp or surgical debridement are likely to be the most appropriate methods for dehisced incisions graded as WUWHS SWD Grades 3/3a and 4/4a.

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Table 11 Main debridement techniques used in dehisced wounds

<table>
<thead>
<tr>
<th>Technique</th>
<th>Description and notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Autolytic</td>
<td>Devitalised tissues are softened and liquefied by enzymes occurring naturally in the wound. Facilitated by dressings that support a moist wound environment. Selective and non-invasive. Can be used before/between other methods of debridement.</td>
</tr>
<tr>
<td>Mechanical</td>
<td>Wet to dry dressings: a moist gauze pad is applied to the wound. As it dries, devitalised tissue becomes attached and is removed with the gauze. Monofilament pad or debridement cloth: devitalised tissue is detached and removed through vigorous cleansing of the wound with the pad or cloth; can be used with autolytic debridement.</td>
</tr>
<tr>
<td>Sharp</td>
<td>Devitalised tissue is removed using a scalpel, scissors and/or forceps. Quick and selective. Requires specialist training; may require local anaesthesia.</td>
</tr>
<tr>
<td>Surgical</td>
<td>Excision of non-viable tissue from wound margins back to viable healthy tissue. Selective. Requires specialist training; may require general anaesthesia and an operating room.</td>
</tr>
<tr>
<td>Larval</td>
<td>Prepared larvae of the green bottle fly (Lucilia sericata) placed in the wound ingest devitalised tissue and bacteria. Selective.</td>
</tr>
<tr>
<td>Ultrasonic</td>
<td>Ultrasound energy is used to break up devitalised tissues; the fragments are washed out with an inbuilt irrigation system. Quick and selective. Requires specialist training.</td>
</tr>
<tr>
<td>Hydrosurgical</td>
<td>A high-pressure jet of saline is used to cut away devitalised tissue. Relatively selective and quick. Requires specialist training.</td>
</tr>
</tbody>
</table>
Cleansing
Wound cleansing aims to remove loose debris, slough, microbes and the remnants of previous dressings from the wound and the surrounding skin\textsuperscript{118}. Cleansing agent selection should be guided by local policy. Cleansing agents include potable water (i.e. water that is safe to drink) or sterile saline\textsuperscript{119,120}.

If the wound is infected, an antimicrobial irrigation solution may be considered for cleansing\textsuperscript{118}. However, the role of antimicrobial irrigation solutions in the management of infected wounds has not yet been fully elucidated.

Management of systemic infection
Patients with systemic signs and symptoms of SSI or erythema extending >5cm from the incision with induration or necrosis should receive a course of systemic antibiotics\textsuperscript{100}. The antibiotics should be selected according to the location of the incision, local antibiotic policy and resistance patterns, and the results of microbiological analysis\textsuperscript{100,121}.

Systemic antibiotics are not usually recommended for the management of a patient with SWD who has only local signs and symptoms of infection. However, this may not apply if the infection is in an incision in which it is important to prevent spreading infection because the consequences may be severe, e.g. a sternomy incision.

Local management of infection
In keeping with guidance on the management of SSI, sutures and clips remaining in a partially dehisced wound should be removed from areas of the incision in which there are signs and symptoms of infection, including abscess\textsuperscript{100}.

Suture/clip removal in a partly dehisced incision should be approached with caution as it may result in expansion or new areas of SWD. Removal should be conducted by a clinician with the appropriate competency and in a care setting that has the facilities to manage the consequences of further dehiscence as appropriate for the location of the incision.

Topical antimicrobials
Topical antimicrobial agents have two main roles in the management of SWD:

- Management of local infection
- Prevention of infection in patients with SWD who are at increased risk of infection.

A wide range of antimicrobial agents is available for use in wounds, including iodine, silver and polyhexamethylene biguanide (PHMB)\textsuperscript{122}. Antimicrobial agents are available in several formulations, including antimicrobial-impregnated dressings (as flat sheets, ribbons or ropes), pastes, gels, powders and irrigation solutions. The properties of an ideal antimicrobial dressing include fast and continued release of the antimicrobial agent into the wound environment to achieve rapid onset and sustained bactericidal activity\textsuperscript{123}. Box 9, page 24, lists the factors that will influence choice of antimicrobial type and formulation.

Two-week challenge
Topical antimicrobials should not be used indefinitely\textsuperscript{93}. Use should be reviewed after two weeks (the ‘2-week challenge’) if monitoring has not indicated that review should take place sooner. If after two weeks the SWD has improved, the antimicrobial should be discontinued. If the SWD has not improved, the patient and the wound should be reassessed and consideration given to changing the topical antimicrobial to a different agent for a further 2-week challenge\textsuperscript{33}.

Topical antimicrobials should be used according to the principles of the ‘2-week challenge’.
Moist wound healing and exudate management

Dressings applied to areas of SWD need to:

- Maintain a moist wound environment to support healing while absorbing excess exudate that could act as a medium for bacterial growth or cause periwound maceration
- Protect the dehisced wound from external contamination and further ingress of microbes.

As discussed above, a dressing may also be used as the delivery vehicle for a topical antimicrobial or as a means of facilitating autolytic debridement.

The dressing selected should be of an absorbency that maintains a moist wound environment without leakage or causing periwound skin damage and that allows for a suitable interval between dressing changes. Ideally, dressing change frequency should tally with the need for wound monitoring: wounds that are infected require more frequent monitoring than uninfected wounds and so are likely to require more frequent dressing changes.

The performance of an individual dressing is affected by the type and quantity of material from which it is constructed. Therefore, it is difficult to make generalisations about the absorbency and exudate handling capability of different dressings. To compound the issue, dressings often combine material types. In very general terms, dressings containing foam, alginate or carboxymethylcellulose are suitable for management of medium to high exudate levels.

The properties of an individual dressing are highly reliant on its construction and constituent materials

The elimination of dead space in deeper SWD is important to prevent accumulation of fluid that may increase risk of infection. Wounds of WUWHS SWD Grades 2 and 3 being treated with dressings will need to be filled with a dressing material (e.g. in rope, ribbon, strip or paste form) appropriate for exudate level. A secondary dressing will be necessary to keep the filler in place. Dehisced wounds of WUWHS SWD Grade 4 are, at least initially, unlikely to be treated with dressings alone.

The dressing(s) selected for an area of dehiscence should be appropriate for the exudate/drainage level and depth of the wound and for the anticipated dressing change frequency

NPWT

Negative pressure wound therapy (NPWT) (Box 10, page 27) is particularly suitable for highly exuding, deep or complex dehisced wounds. NPWT fulfils the needs for moist wound healing, exudate/drainage management, elimination of dead space and protection from external contamination in the facilitation of healing by secondary or tertiary healing after SWD.

NPWT has been used for the management of a wide range of acute and chronic wound types for more than 20 years. The extent of the evidence and clinical experience behind this treatment modality has resulted in NPWT being suggested as the ‘gold standard treatment’ for open abdominal wounds and dehisced sternal wounds.

The considerable number of studies that have investigated the use of NPWT on infected surgical wounds often include mixed populations of patients with and without SWD. Reports focusing largely on the role of NPWT in SWD include for:

- Abdominal wound dehiscence
- Post-sternotomy dehiscence
- Post-caesarean dehiscence
- Perineal dehiscence
- Dehiscence after amputation
More recently, the use of NPWT over closed surgical incisions has been shown to reduce rates of SSI, seroma/haematoma and dehiscence, and to improve scar quality\(^{140,141}\).

**Mode of action**

NPWT applies controlled negative pressure to a wound or incision, provides a physical barrier to external contamination and removes excess wound drainage. In addition, in open wounds NPWT aids healing by:

- Contracting wound edges to reduce wound size
- Stimulating angiogenesis and granulation tissue formation
- Reducing oedema
- Improving tissue perfusion\(^{142-145}\) (Figure 14).

**Box 10** Negative pressure wound therapy (NPWT) in open wounds

- NPWT involves the application of controlled negative pressure (suction) over an open wound (or closed surgical incision\(^*\)) and perilesional tissues
- A wound filler, e.g. foam or gauze, and sometimes a liner, is placed in the wound and an adhesive film is used to cover the wound and form a seal
- The seal allows delivery of suction generated by an electrically-powered pump (that contains batteries or is plugged into a mains electricity source) or by a mechanically-powered pump
- NPWT devices vary in size, portability and format, e.g. some include a canister of varying volume for collection of fluids while others employ absorption and evaporation for fluid handling; some are designed for single-use
- Some single-use NPWT (sNPWT) devices that use the dressing for fluid management and as a wound interface allow wider coverage of the periwound area
- Some NPWT devices for use in open wounds incorporate instillation of solutions such as normal saline or antimicrobials

*For information on the mode of action of NPWT in closed surgical incisions, see page 36

**Figure 14** Mode of action of NPWT in open wounds

**Role of NPWT in the management of SWD**

NPWT has several potential roles in the management of SWD, e.g. following primary closure of the dehisced wound, during healing by secondary intention and in preparation for delayed primary closure (Figure 15, page 28 and Figure 16, page 29). The type of NPWT device selected is dependent on several factors (Box 11, page 28).

NPWT has been widely used in the management of SWD and is increasingly being used to prevent SWD.

NPWT should be used in the context of appropriate wound bed preparation (debridement) and management of infection, if present.

Risk of delayed healing (Box 4, page 7) may be an indication for the use of NPWT in the management of patients with SWD.

With the increasing use of closed incision NPWT for prophylaxis of surgical site complications, some patients with SWD may previously have received this treatment modality. The potential benefits and harms of using NPWT again on previously treated dehisced wounds are not yet known, and some clinicians would approach reuse of NPWT with caution.
**NPWT and the management of infection**

Recent recommendations on the use of NPWT state that NPWT should be used only as an adjunctive treatment in the management of wound infection\(^{142}\). The same recommendations comment that the use of antimicrobial dressings or fillers under NPWT, e.g. PHMB-impregnated gauze or silver-impregnated foam, may aid infection control\(^{142}\).

Clinicians should check the indications, contraindications and cautions for the specific NPWT device under consideration.

**Box 11 | Factors involved in selecting the type of NPWT for use in the management of SWD**

- **Contraindications and cautions** (Box 12, page 29) – clinicians should always consult the manufacturer’s instructions for the NPWT device under consideration before implementing use in a patient.

- **Location of the incision/dehiscence** – the dressing needs to be able to conform to the three-dimensional shape of the anatomical area sufficiently well to avoid dead space and to form the seal needed for NPWT to work.

- **Volume of wound drainage** – the device selected should be able to cope with the anticipated volume of drainage, e.g. if wound drainage is <300ml/week canister-less single-use NPWT (sNPWT) may be appropriate; if drainage is >300ml/week a canister-based device of appropriate capacity will be more suitable.

- **Depth of the wound** – e.g. some sNPWT devices should be used on wounds with a maximum depth of 2cm; some sNPWT devices cannot be used with fillers.

- **Size (area) of the wound** – the NPWT device selected must be appropriate for the size (area) of the wound.

- **Infection** – an antimicrobial interface may be required and should be compatible with the NPWT device being used; if NPWT with instillation is selected, the device needs to be instillation-capable.

- **Care setting** – the NPWT device selected should be of a type that can be cared for appropriately and safely in the setting in which it will be used.

- **Patient needs and acceptance** – e.g. patients who are active or able to return to work are likely to prefer a portable sNPWT device.

**Figure 15 | Potential roles of NPWT in the management of SWD**

NPWT should be used in the context of holistic management of the patient (see Figures 10 and 11, pages 21 and 22) and take into account the contraindications/cautions for the NPWT device being considered.

- Consider closed incision prophylactic NPWT if patient has risk factors for surgical site complications (e.g. Box 4, page 7)

- Consider NPWT if wound drainage is requiring very frequent dressing changes and/or if the patient has risk factors for delayed healing (e.g. Box 4, page 7); fillers may be required for deep wounds

- Consider NPWT + antimicrobial interface if the wound is at high risk of infection

- Consider NPWT + antimicrobial interface for infected wounds, or for larger infected wounds, consider NPWT + instillation/irrigation

- Consider NPWT with fillers if wound drainage is requiring very frequent dressing changes and/or if the patient has risk factors for delayed healing (e.g. Box 4, page 7)

- Consider NPWT + antimicrobial interface if the wound is at high risk of infection

- Consider NPWT + antimicrobial interface for infected wounds, or for larger infected wounds, consider NPWT + instillation/irrigation

- NPWT may also be indicated after primary closure or after closure with a flap or graft.
Necrotic tissue with eschar
Osteomyelitis
Non-enteric and unexplored fistulae
Malignancy in the wound – unless part of palliative care
Exposed blood vessels
Exposed nerves
Exposed anastomotic site
Exposed organs
 Patients at high risk for bleeding – e.g. from a blood clotting disorder
 Patients on anticoagulants or platelet aggregation inhibitors
 Patients with:
- Friable and infected blood vessels
- Vascular anastomosis
- Treated infected wounds
- Sharp edges in the wound – e.g. bone fragments
- Spinal cord injury
- Enteric fistulae
Patients requiring:
- Magnetic resonance imaging (MRI)
- Treatment in a hyperbaric oxygen chamber (HBOT)
- Defibrillation
- Use near the vagus nerve (may cause bradycardia)

N.B. The information in this box is a generalised list of contraindications and cautions to the use of NPWT. Clinicians should check the contraindications and cautions for the specific NPWT device under consideration

- Pre-application
- Single-use NPWT applied

**NPWT with instillation**

NPWT with instillation has been developed to allow the delivery of topical solutions, such as saline and antimicrobial agents, to the wound bed while maintaining a seal over the wound. During the periodic introduction of the solution to the wound bed, the vacuum pump is halted for a short time, e.g. 20 minutes, and then restarted until the next episode of instillation.

NPWT with instillation may be used in the management of infection in acute and chronic wounds because it reduces bioburden. Much remains to be learnt about which instillation solutions to use, and when and for
how long, but a review of current evidence suggests that larger (area >40cm²) or deeper wounds and wounds that have high bacterial bioburden may be the most appropriate indications.546

Changing NPWT modality
With the range of NPWT devices available there is scope for patients to be moved from one device to another as treatment progresses and therapeutic requirements change, e.g. as wound size, depth and/or exudate level decrease, the patient is discharged if in hospital, the patient becomes increasingly mobile and/or returns to work. The need for an alternative NPWT device should be assessed on an individual basis and with reference to local policy/wound care formulary where available. If appropriate, advice should be taken from tissue viability or medical teams.

NPWT in the community
A UK survey found that over half of patients with surgical wounds healing by secondary intention were cared for in community settings.53 Because of the trend for decreasing length of hospital inpatient stays and the development of portable NPWT devices, clinicians working in the community are increasingly likely to be involved in the care of patients with SWD who have been discharged with NPWT or who have commenced NPWT post-discharge.547

Clinicians in the community play important roles in supporting, monitoring and managing patients receiving treatment with NPWT to ensure it is used safely, appropriately and effectively. Such involvement may involve liaising with clinicians in other services and managing transitions between NPWT devices and other methods of wound management as needed.

Patients being treated in community settings need information, education and training, as appropriate, about their NPWT device and how to use it. They also need to know how to contact a clinician and the general and SWD-related signs and symptoms and issues with the NPWT device should trigger contact.548

Control of oedema
Inflammation, an integral part of the healing response following surgery, increases permeability of blood vessels and causes interstitial fluid accumulation that may manifest clinically as oedema. Post-operative swelling due to oedema may particularly be a problem in lower limb surgery, e.g. following ankle surgery or saphenous vein harvesting, and may contribute to SWD because it delays healing.29

Control of oedema in patients with lower limb SWD may aid healing
Strategies to reduce oedema include limb elevation and/or the use of compression therapy, e.g. bandages, compression stockings or intermittent pneumatic compression.55 The ankle-brachial pressure index (ABPI) of patients being considered for lower limb compression therapy should be ascertained to assess arterial blood supply.55

Monitoring and reassessment
Patients with SWD should be monitored carefully at each dressing or device change, including for signs and symptoms of infection. Management should be adjusted as indicated by the reassessment and if necessary referrals made to a tissue viability or surgical service.

A full reassessment of the dehisced wound and current management should take place at two weeks for infected SWD and at four weeks for uninfected SWD unless monitoring indicates the need for full reassessment sooner.
60-year-old woman
WUWHS SWD Grade 1a of the dermal layer that affected a 2cm section of an otherwise healed incision following laparotomy 10 days previously
Draining pus; no signs of systemic infection
Local wound infection resolved after treatment for one week with a topical antimicrobial (silver) dressing
After discontinuation of the silver dressing, a foam dressing was applied
Wound was fully healed within 3 weeks of presentation

45-year-old woman
WUWHS SWD Grade 2 of the dermal layer and subcutaneous layers affecting over 50% of an incision made 8 days previously for removal of a non-malignant breast lump
No clinical signs or symptoms of infection
Wound was packed with an alginate dressing and a secondary foam dressing was applied
Wound was fully healed 2 weeks later

58-year-old man
WUWHS Grade 3 SWD affecting the full length of the incision with separation of the full thickness of the skin and subcutaneous tissue and fascial exposure, following spinal surgery 3.5 weeks previously
Wound was clean and not infected
Plastic surgery team was consulted and the patient returned to the operating room, where he underwent tissue undermining and paraspinous muscle mobilisation, layered tissue closure, drain insertion and NPWT over the closed incision
Wound was fully healed within 2 weeks of presentation

62-year-old man
WUWHS Grade 3a SWD of approximately 50% of the incision with separation of the skin and subcutaneous tissue and fascial exposure, following spinal surgery 5 weeks previously
He had developed haematoma and dehiscence after discharge home. His re-presentation was delayed because he lived a considerable distance from the hospital
Wound contained pus; there were no signs of systemic infection
Local wound care was performed with antimicrobial wound dressing changes followed by wound closure with trapezius flap
Wound was fully healed within 6 weeks of presentation

70-year-old woman
WUWHS SWD Grade 4 mechanical dehiscence of a sternal incision that extended to sternal bone following CABG 6 days previously
Minimal serous exudate; no signs of local or systemic infection
Had decided not to wear her bra post-operatively, which would have offered support to the incision
NWPT was commenced with foam filler (no liner) at -120mmHg; discontinued 10 days later
Wound was then managed with a carboxymethylcellulose and foam adhesive dressing until fully healed at 3 weeks
Patient was advised to wear her bra and not to undertake any heavy lifting or pulling/pushing for up to 12 weeks post-healing
The large number of risk factors associated with SWD (Table 6, page 10, and Table 7, page 11) provide multiple opportunities before, during and after surgery to implement interventions that aim to reduce risk.

Keys to prevention of SWD are identifying patients at risk, modifying risk of SWD and SSI where possible, implementing preventative measures, and post-operative monitoring for healing progress and signs of infection or possible dehiscence (Figure 17).

Risk assessment

In elective surgery, pre-operative consultations provide opportunities for thorough risk assessment. The risk assessment can then be used to explain to a patient their individual level of risk for SWD and other post-operative complications, and to plan risk reduction for patient-related modifiable risk factors (such as high BMI and smoking).

In emergency surgery, risk assessment also has an important role. However, opportunities for discussion of risk levels and amelioration of patient-related modifiable risk factors may be limited.

Risk for SWD should be assessed pre-operatively and taken into account when planning surgery. Depending on the indication for surgery, if risk of SWD is high, thought may need to be given to whether surgery remains appropriate.

Calculators of risk for SWD

In practice, risk for SWD is often ascertained pre-operatively by clinical observation. However, risk calculators can be used to provide an objective assessment of risk.

Risk calculators specific to SWD

Two scoring systems have been developed and validated for the evaluation of risk for SWD in patients undergoing laparotomy: the Veterans Affairs Medical Center (VAMC) and Rotterdam risk models (Appendix 3, page 40). Both models include post-operative variables, and the VAMC model includes intra-operative variables, suggesting there may be limitations in using these models for pre-operative risk assessment for SWD. A further unvalidated scoring system of risk for SWD has also been developed (Appendix 3, page 40).

A comparison of the VAMC and Rotterdam risk models concluded that both can be used to predict abdominal SWD. A further study of the Rotterdam risk model reported that the global risk score (i.e. the score using all variables) had better accuracy than the pre-operative risk score (i.e. the score that excluded the post-operative variables).
The Breast Reconstruction Risk Assessment (BRA) tool (www.brascore.org) uses a range of patient-related factors including height, weight, age, chemotherapy, comorbidities and bleeding risks to calculate risk for a range of surgical complications for a range of reconstructive modalities. Risks calculated include for dehiscence, SSI and seroma.

Assessment of risk for SSI

SSI is a major risk factor for SWD. Tools that indicate increased risk for SSI (Box 13) may therefore indicate increased risk for SWD.

The outcome of pre-operative risk assessment for SWD and other post-operative complications along with the specific risk factors identified should be clearly documented and communicated to all members of the team caring for the patient before, during and after surgery.

Reducing risk of SWD

Reducing risk of SWD includes pre-operative modification of comorbidities and optimisation of patient condition, excellent surgical technique, selection of the appropriate closure method, oedema prevention or reduction, minimisation of SSI risk, post-operative monitoring and patient education.

Comorbidity risk modification

Patients should be referred as appropriate for pre-operative risk modification, e.g. weight loss and smoking cessation programmes, improved control of diabetes mellitus, nutrition management. In emergency surgery, however, opportunities to influence modifiable risk factors will be more limited. In some cases, it may be appropriate to delay surgery to reduce the risk of SWD, e.g. to allow more time for the patient to lose weight and cease smoking or to recover from radiation therapy.

Surgical technique

Excellent surgical technique is likely to lessen the risk of SWD by reducing problems with healing, decreasing haematoma and seroma formation, and lessening the risk of SSI. Examples of excellent technique include gentle handling of tissues, meticulous control of bleeding, maintenance of blood supply, prevention of tissue drying, removal of devitalised or contaminated tissues, avoidance of dead space, and the use of an appropriate closure technique.

The wound closure technique selected for primary closure should be appropriate for the site of the incision and surgical procedure, and should ensure that the tissue layers are accurately apposed and tension across the incision is minimised. Minimising tension may require suturing of individual tissue layers and careful consideration of the spacing and length of the sutures.

For some patients, primary closure of the incision is not appropriate because of increased risk of SWD or other complications (e.g. infection, haemorrhage or abdominal compartment syndrome). In such cases, the incision may be left open (with an appropriate protective covering/device) until a time when closure is appropriate or possible.

Delayed primary closure of the initial incision may be used to avoid dehiscence in patients recognised to be at increased risk of SWD.
**Oedema reduction/prevention**

Oedema may contribute to SWD because it may hinder healing by impairing tissue perfusion and increasing tension in the incision because of tissue swelling.

**Gentle tissue handling during surgery, careful fluid management and treatment of infection may reduce the risk of SWD by decreasing oedema formation**

Local cooling of the incision, e.g. through the application of icepacks (cryotherapy), is often used to reduce pain following orthopaedic surgery, but may also reduce oedema. Compression may also help to reduce oedema formation and has been reported to reduce surgical wound complications following total ankle arthroplasty\(^{161}\). Combinations of cryotherapy and compression may also be used\(^{162}\).

**The ankle–brachial pressure index (ABPI) of a patient being considered for lower limb compression following surgery should be ascertained to exclude arterial insufficiency**

**Incision management**

Epithelialisation of surgical incisions is usually complete, i.e. the wound is usually sealed, within 48 hours of surgery. Therefore, dressings applied to an incision are usually left in place for at least the first 48 hours post-operatively while being inspected regularly\(^{163}\). The ideal post-operative dressing acts as a barrier to bacteria, is vapour-permeable (i.e. allows water to evaporate), allows monitoring of fluid accumulation, and has a low risk of causing trauma or blistering\(^{164}\).

The World Health Organization (WHO) guideline on the prevention of SSI has made a conditional recommendation regarding NPWT: “The panel suggests the use of prophylactic negative pressure wound therapy (pNPWT) in adult patients with primarily closed surgical incisions in high-risk wounds, for the purpose of the prevention of SSI...”\(^{163}\). NPWT on closed incisions has been reported to also decrease the incidence of SWD\(^{141,165–168}\). In common with many other wound products, research into NPWT is ongoing. The protocols for several randomised controlled studies have been published\(^{10,169–171}\) or are available at: clinicaltrials.gov.

**Overview of interventions to reduce SWD and SSI**

Several national and international guidelines aimed at reducing the occurrence of SSI have been developed\(^{121,163,172,177}\). As there is overlap between the risk factors for SWD and those for SSI, and SSI can cause SWD, the interventions recommended in the guidelines on SSI prevention also have relevance to the prevention of SWD.

Table 12, page 35, and Appendix 4, pages 41–42, list interventions aimed at reducing risk of surgical site complications such as SWD and SSI arranged according to the phase (planning, pre-operative, intra-operative and post-operative) of surgery to which they relate.

The use of interventions to reduce the risk of SWD and SSI should take place in the context of a full assessment of the patient and the implementation of other safety interventions, e.g. prevention of deep vein thrombosis (DVT) and pressure ulcers (PUs).
### Table 12 | Interventions for reduction of risk of surgical site complications such as SWD and SSI

See Appendix 4, pages 41–42, for more detail.

<table>
<thead>
<tr>
<th>Phase</th>
<th>Intervention</th>
<th>Planning</th>
<th>Pre-operative</th>
<th>Intra-operative</th>
<th>Post-operative</th>
</tr>
</thead>
<tbody>
<tr>
<td>Planning</td>
<td>Education of patient/carer/family and management of expectations</td>
<td>✔</td>
<td>✔</td>
<td>✔</td>
<td>✔</td>
</tr>
<tr>
<td>Planning</td>
<td>Assessment and management/optimisation of comorbidities - e.g. obesity, malnutrition, diabetes mellitus, COPD, anaemia, cardiovascular disease</td>
<td>✔</td>
<td>✔</td>
<td>✔</td>
<td>✔</td>
</tr>
<tr>
<td>Planning</td>
<td>Screening for nasal carriage of <em>Staphylococcus aureus</em> and decolonisation according to local protocol – e.g. test patients undergoing cardiac surgery or surgery involving an implant (e.g. arthroplasty or breast implant) and those who are healthcare workers or institutional residents</td>
<td>✔</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Planning</td>
<td>Management of bleeding/thrombotic risk in patients on oral anticoagulants</td>
<td>✔</td>
<td>✔</td>
<td>✔</td>
<td>✔</td>
</tr>
<tr>
<td>Planning</td>
<td>Consider nutritional supplementation</td>
<td>✔</td>
<td>✔</td>
<td>✔</td>
<td>✔</td>
</tr>
<tr>
<td>Pre-operative</td>
<td>Use of an operative safety checklist – e.g. WHO Surgical Safety Checklist</td>
<td>✔</td>
<td>✔</td>
<td>✔</td>
<td>✔</td>
</tr>
<tr>
<td>Pre-operative</td>
<td>Maintenance of normothermia, unless otherwise indicated</td>
<td>✔</td>
<td>✔</td>
<td>✔</td>
<td>✔</td>
</tr>
<tr>
<td>Pre-operative</td>
<td>Monitor and control blood glucose of patients with diabetes mellitus</td>
<td>✔</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pre-operative</td>
<td>Showering or bathing by patient on day of surgery using plain or antimicrobial soap/cleanser</td>
<td>✔</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pre-operative</td>
<td>Use of clippers (rather than a razor) for hair removal</td>
<td>✔</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pre-operative</td>
<td>Location of heparin injection sites away from operative site</td>
<td>✔</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pre-operative</td>
<td>Management of hydration/fluid levels to produce normovolaemia, while avoiding fluid overload and hypovolaemia</td>
<td>✔</td>
<td>✔</td>
<td>✔</td>
<td>✔</td>
</tr>
<tr>
<td>Pre-operative</td>
<td>Maintenance of adequate tissue perfusion</td>
<td>✔</td>
<td>✔</td>
<td>✔</td>
<td>✔</td>
</tr>
<tr>
<td>Pre-operative</td>
<td>Timely administration of prophylactic antibiotics as indicated by local guidelines</td>
<td>✔</td>
<td>✔</td>
<td>✔</td>
<td>✔</td>
</tr>
<tr>
<td>Pre-operative</td>
<td>Administration of antifibrinolytic agents as indicated by local guidelines to reduce blood loss and need for blood transfusion</td>
<td>✔</td>
<td>✔</td>
<td>✔</td>
<td>✔</td>
</tr>
<tr>
<td>Intra-operative</td>
<td>Compliance with hygiene measures by operating room personnel</td>
<td>✔</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Intra-operative</td>
<td>Minimisation of operating room traffic</td>
<td>✔</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Intra-operative</td>
<td>Optimal oxygenation</td>
<td>✔</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Intra-operative</td>
<td>Skin preparation with an antiseptic immediately prior to incision</td>
<td>✔</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Intra-operative</td>
<td>Use of an iodophor-impregnated drape, unless the patient has an iodine allergy, if an incise drape is necessary</td>
<td>✔</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Intra-operative</td>
<td>Use of excellent surgical technique with gentle handling of tissues, meticulous control of bleeding and avoidance of dead space</td>
<td>✔</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Intra-operative</td>
<td>Avoidance of tension across incision</td>
<td>✔</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Intra-operative</td>
<td>Use of wound edge protectors/guards during laparotomy</td>
<td>✔</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Intra-operative</td>
<td>Intra-operative wound irrigation</td>
<td>✔</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Intra-operative</td>
<td>Change of gloves during procedure and/or before closure of wound; double gloving</td>
<td>✔</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Intra-operative</td>
<td>Senior/experienced surgeon performing closure</td>
<td>✔</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Intra-operative</td>
<td>Use of gentamicin-impregnated collagen sponges</td>
<td>✔</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Intra-operative</td>
<td>Use of triclosan-coated sutures</td>
<td>✔</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Intra-operative</td>
<td>Covering of the incision(s) with a dry absorbent sterile dressing under sterile conditions and before the patient leaves the operating room</td>
<td>✔</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Intra-operative</td>
<td>Consider prophylactic NPWT (e.g. single-use NPWT) for patients at increased risk of SSI or SWD</td>
<td>✔</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Post-operative</td>
<td>Maintenance of the dressing over the incision for at least 48 hours unless there are signs and symptoms indicating earlier inspection is warranted</td>
<td>✔</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Post-operative</td>
<td>Cryotherapy (e.g. application of ice) and compression</td>
<td>✔</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Post-operative</td>
<td>Visitor restrictions and hygiene measures – e.g. hand cleansing/protective clothing as appropriate</td>
<td>✔</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Post-operative</td>
<td>Monitor incision for healing progress and signs/symptoms of dehiscence or infection</td>
<td>✔</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Post-operative</td>
<td>Patient Reported Outcome/Experience Measures (PROMS/PREMS) or questionnaires</td>
<td>✔</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Post-operative</td>
<td>Perform surveillance of post-operative wound complications and compliance with surgical wound complication reduction bundles</td>
<td>✔</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Prevention of SWD with prophylactic NPWT

In addition to roles in the management of surgical incisions healing by secondary intention or being managed with delayed primary closure, there is established and growing evidence that prophylactic NPWT, including single-use NPWT (sNPWT), reduces the incidence of surgical site complications, including SWD and SSI.120,129,174 A recent study of patients undergoing routine primary hip and knee replacements found that use of prophylactic sNPWT produced cost-savings in an analysis of all patients, with greater savings in subgroups of patients at higher risk of surgical site complications.175

Mode of action of prophylactic NPWT on closed surgical incisions

In open wounds, NPWT has been found to have effects that may be relevant to closed incisions, e.g. stimulation of angiogenesis and reduction of oedema.176,177 In addition to aiding exudate management and protecting the incision from external contamination, prophylactic NPWT used in the management of closed surgical incisions has also been shown to:

- Reduce lateral tension (Box 14)
- Improve lymphatic clearance
- Reduce seroma and haematoma formation.140

Prophylactic NPWT may also have effects in the tissues surrounding the incision (the ‘zone of injury’) by reducing oedema and levels of inflammatory markers and may promote collagen synthesis.183 Together these effects may contribute to faster and stronger healing, and reduced risk of SWD.20

Effect of prophylactic NPWT on rates of dehiscence

Individual studies of prophylactic NPWT, including studies of sNPWT, in orthopaedic and breast surgery and a recent meta-analysis of effect in a range of surgery types have found significant reductions in rates of SWD (Table 13, page 37). However, other published systematic reviews and meta-analyses have found that study heterogeneity prevented analysis or that the evidence for reductions in SWD is inconclusive.35,140,184–187 Protocols for ongoing trials into the effect of prophylactic NPWT on rates of SWD have been published.10,171

Selecting patients for prophylactic NPWT

Figure 18, page 37, proposes a role for prophylactic NPWT in the prevention of SWD in patients likely to be at increased risk. It is an adaptation of the proposed role of NPWT for the prevention of surgical site complications in closed surgical incisions that appears in a recent international consensus document.20

The prophylactic NPWT device selected will depend on factors including the location and size of the closed surgical incision, the anticipated level of drainage from the incision, and the other needs of the patient. For example, a canister-less prophylactic sNPWT device may be selected for a patient who has a closed surgical incision that is likely to have low levels of drainage and who is able to regain mobility and return to work soon after surgery.

N.B. The procedures given here are examples and do not comprise a complete list of procedures which have a high rate of surgical wound complications that could have severe consequences, e.g. failure of surgery, life-changing implications and death. Individual patients undergoing the same procedure may experience different levels of risk and severity of consequences of surgical site complications as a result of variation in the presence of other risk factors.

Box 14 | Effects of prophylactic NPWT on stresses in closed incisions

- During computer modelling, prophylactic NPWT reduced the lateral tension inherent in the incision by 45%–70%.179,180
- About 50% more force was required in a physical model to disrupt an incision to which prophylactic NPWT was applied than to disrupt an incision closed with sutures or clips.179,180
- Prophylactic NPWT increased the breaking strength of wounds in animal studies.181–183

Box 15 | Examples of higher consequence/higher incidence procedures for surgical site complications

- Complex surgery – e.g. major colorectal surgery, oesophagogastrectomy, extensive combined procedures which include a long skin-to-skin time, especially in redo or multiple redo procedures
- Arthroplasty revision
- Surgery involving high energy below knee fractures
- Major oncological procedures in children
- After radiotherapy

N.B. The procedures given here are examples and do not comprise a complete list of procedures which have a high rate of surgical wound complications that could have severe consequences, e.g. failure of surgery, life-changing implications and death. Individual patients undergoing the same procedure may experience different levels of risk and severity of consequences of surgical site complications as a result of variation in the presence of other risk factors.

Post-operative patient education

Post-operatively, patients should be advised on appropriate levels of activity, dressing/device care, signs and symptoms of SWD and SSI, and when and who to contact with problems.
### Table 13 | Reductions in rates of SWD in studies of prophylactic NPWT on closed surgical incisions

<table>
<thead>
<tr>
<th>Author &amp; Year</th>
<th>Type(s) of surgery</th>
<th>Details</th>
<th>SWD rates</th>
</tr>
</thead>
<tbody>
<tr>
<td>Strugala &amp; Martin, 2017</td>
<td>Mixed: breast, orthopaedic, caesarean section, coronary artery bypass graft</td>
<td>Meta-analysis of 6 studies; 1068 patients; 1291 incisions</td>
<td>n=611 NPWT*; n=680 control SWD: 12.8% vs 17.4% (p&lt;0.05)</td>
</tr>
<tr>
<td>Stannard et al, 2012</td>
<td>Orthopaedic (lower limb)</td>
<td>249 patients; 263 fractures</td>
<td>n=141 NPWT**; n=122 control SWD: 8.6% vs 16.5% (p&lt;0.05)</td>
</tr>
<tr>
<td>Galiano et al, 2014</td>
<td>Breast</td>
<td>199 patients; 398 incisions</td>
<td>n=199 NPWT*; n=199 control SWD at 21 days: 16.2% vs 26.4% (p&lt;0.05)</td>
</tr>
<tr>
<td>Adogwa et al, 2014</td>
<td>Orthopaedic (spine)</td>
<td>160 patients</td>
<td>n=46 NPWT**; n=114 control SWD: 6.38% vs 12.28% (p&lt;0.05)</td>
</tr>
<tr>
<td>Holt &amp; Murphy, 2015</td>
<td>Therapeutic mammoplasty and symmetrising reduction</td>
<td>24 patients</td>
<td>n=24 NPWT*; n=24 control SWD: 4.2% vs 16.7% (p not reported)</td>
</tr>
</tbody>
</table>

*PICO Single Use Negative Pressure Wound Therapy (Smith & Nephew); ** V.A.C. (KCI/Acelity) |
CS: case series; MA: meta-analysis; RCT: randomised controlled trial; RS: retrospective study

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**RESEARCH NEEDS**

Evidence is accumulating of the large scale of surgical wound healing problems and the high economic and social costs they bring to healthcare systems and patients. More research is needed to further clarify the health economic impact of SWD, including incidence (associated and not associated with infection), quality of life data, costs of management in hospital and community settings and the impact of interventions to prevention SWD.
## Appendix 1. Centers for Disease Control and Prevention (CDC) definitions of SSI

<table>
<thead>
<tr>
<th>Type of SSI</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Superficial incisional SSI</strong>*</td>
<td>Infection occurs within 30 days after any operative procedure (where day 1 = the procedure date) AND involves only skin and subcutaneous tissue of the incision AND the patient has at least one of the following: a. purulent drainage from the superficial incision b. organisms identified from an aseptically-obtained specimen from the superficial incision or subcutaneous tissue by a culture- or non-culture-based microbiologic testing method which is performed for purposes of clinical diagnosis or treatment (e.g. not Active Surveillance Culture/Testing (ASC/AST)) c. superficial incision that is deliberately opened by a surgeon, attending physician or other designee and culture or non-culture based testing is not performed AND The patient has at least one of the following signs or symptoms: pain or tenderness; localized swelling; erythema; or heat. d. diagnosis of a superficial incisional SSI by the surgeon or attending physician or other designee</td>
</tr>
<tr>
<td><strong>Deep incisional SSI</strong>*</td>
<td>Infection occurs within 30 or 90 days** after the procedure (where day 1 = the procedure date) AND involves deep soft tissues of the incision (e.g. fascial and muscle layers) AND the patient has at least one of the following: a. purulent drainage from the deep incision b. a deep incision that spontaneously dehisces, or is deliberately opened or aspirated by a surgeon, attending physician or other designee and organism is identified by a culture or non-culture based microbiologic testing method which is performed for purposes of clinical diagnosis or treatment (e.g. not Active Surveillance Culture/Testing (ASC/AST)) or culture or non-culture based microbiologic testing method is not performed AND The patient has at least one of the following signs or symptoms: fever (&gt;38°C); localized pain or tenderness. A culture or non-culture based test that has a negative finding does not meet this criterion c. an abscess or other evidence of infection involving the deep incision that is detected on gross anatomical or histopathologic examination, or imaging test</td>
</tr>
<tr>
<td><strong>Organ/space SSI</strong>**</td>
<td>Infection occurs within 30 or 90 days** after the procedure (where day 1 = the procedure date) AND infection involves any part of the body deeper than the fascial/muscle layers, that is opened or manipulated during the operative procedure AND The patient has at least one of the following: a. purulent drainage from a drain that is placed into the organ/space (e.g. closed suction drainage system, open drain, T-tube drain, CT guided drainage) b. organisms are identified from an aseptically-obtained fluid or tissue in the organ/space by a culture or non-culture based microbiologic testing method which is performed for purposes of clinical diagnosis or treatment (e.g. not Active Surveillance Culture/Testing (ASC/AST)) c. an abscess or other evidence of infection involving the organ/space that is detected on gross anatomical or histopathologic examination, or imaging test evidence suggestive of infection</td>
</tr>
</tbody>
</table>

*Superficial incisional SSI and deep incisional SSI may both be further categorised as primary or secondary according to whether the incision in question is the primary incision or is the secondary incision in an operation with more than one incision

**90-day surveillance is for breast surgery, cardiac surgery, coronary artery bypass graft with both chest and donor site incisions, coronary artery bypass graft with chest incision only, craniotomy, spinal fusion, open reduction of fracture, hemorhaphy, hip prosthesis, knee prosthesis, pacemaker surgery, peripheral vascular bypass surgery, ventricular shunt. Some SSI classifications do not specify length of surveillance according to procedure type other than to specify 30 days if no implant is in place, or within one year if an implant is in place.***

**Some SSI classifications based on the CDC classification include diagnosis of organ/space SSI by a surgeon or physician.***

### Public Health England (PHE) definitions of SSI

The definitions of SSI used by Public Health England are based on those established by the US Centers for Disease Control and Prevention (CDC) with minor modifications:
- A requirement for pus cells in addition to a positive culture from wound samples (for all SSI types)
- The need for at least two symptoms to accompany a clinical diagnosis (superficial SSIs only)
- Timing is appearance of SSI within 30 days for all procedures, unless an implant is in place when it is one year.
## Appendix 2. ASEPSIS grading system

<table>
<thead>
<tr>
<th>Criterion</th>
<th>Points</th>
<th>Points</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>10</td>
<td>5 10</td>
</tr>
<tr>
<td>S</td>
<td></td>
<td></td>
</tr>
<tr>
<td>E</td>
<td></td>
<td></td>
</tr>
<tr>
<td>P</td>
<td></td>
<td></td>
</tr>
<tr>
<td>S</td>
<td></td>
<td></td>
</tr>
<tr>
<td>I</td>
<td></td>
<td></td>
</tr>
<tr>
<td>S</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*Scoring is according to proportion of wound affected:*

<table>
<thead>
<tr>
<th></th>
<th>0%</th>
<th>&lt;20%</th>
<th>20%-39%</th>
<th>40%-59%</th>
<th>60%-79%</th>
<th>≥80%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Serous discharge</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
<td>5</td>
</tr>
<tr>
<td>Erythema</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
<td>5</td>
</tr>
<tr>
<td>Purulent exudate</td>
<td>0</td>
<td>2</td>
<td>4</td>
<td>6</td>
<td>8</td>
<td>10</td>
</tr>
<tr>
<td>Separation of</td>
<td>0</td>
<td>2</td>
<td>4</td>
<td>6</td>
<td>8</td>
<td>10</td>
</tr>
<tr>
<td>deep tissues</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Category of infection</th>
</tr>
</thead>
<tbody>
<tr>
<td>ASEPSIS score</td>
</tr>
<tr>
<td>0-10</td>
</tr>
<tr>
<td>11-20</td>
</tr>
<tr>
<td>20-30</td>
</tr>
<tr>
<td>31-40</td>
</tr>
<tr>
<td>&gt;40</td>
</tr>
</tbody>
</table>
### Appendix 3. Veterans Affairs Medical Center (VAMC), Rotterdam and Mir risk scoring systems for abdominal SWD following laparotomy

<table>
<thead>
<tr>
<th>Variables and scores</th>
<th>VAMC&lt;sup&gt;44&lt;/sup&gt;</th>
<th>Rotterdam&lt;sup&gt;45&lt;/sup&gt;</th>
<th>Mir et al, 2016&lt;sup&gt;46&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>Variable</td>
<td>Score</td>
<td>Variable</td>
<td>Score</td>
</tr>
<tr>
<td>CVA with no residual deficit</td>
<td>4</td>
<td>40-49</td>
<td>0.4</td>
</tr>
<tr>
<td>History of COPD</td>
<td>4</td>
<td>50-59</td>
<td>0.9</td>
</tr>
<tr>
<td>Current pneumonia</td>
<td>4</td>
<td>60-69</td>
<td>0.9</td>
</tr>
<tr>
<td>Emergency procedure</td>
<td>6</td>
<td>&gt;70</td>
<td>1.1</td>
</tr>
<tr>
<td>Operative time &gt;2.5 hours&lt;sup&gt;*&lt;/sup&gt;</td>
<td>2</td>
<td>Male gender</td>
<td>0.7</td>
</tr>
<tr>
<td>PGY 4 resident as surgeon&lt;sup&gt;*&lt;/sup&gt;</td>
<td>3</td>
<td>Chronic pulmonary disease</td>
<td>0.7</td>
</tr>
<tr>
<td>Clean wound classification&lt;sup&gt;*&lt;/sup&gt;</td>
<td>-3</td>
<td>Ascites</td>
<td>1.5</td>
</tr>
<tr>
<td>Superficial wound infection&lt;sup&gt;*&lt;/sup&gt;</td>
<td>5</td>
<td>Jaundice</td>
<td>0.5</td>
</tr>
<tr>
<td>Deep wound infection&lt;sup&gt;*&lt;/sup&gt;</td>
<td>17</td>
<td>Anaemia</td>
<td>0.7</td>
</tr>
<tr>
<td>Failure to wean from ventilator&lt;sup&gt;*&lt;/sup&gt;</td>
<td>6</td>
<td>Emergency surgery</td>
<td>0.6</td>
</tr>
<tr>
<td>One or more complications other than dehiscence&lt;sup&gt;**&lt;/sup&gt;</td>
<td>7</td>
<td>Gallbladder/bile duct</td>
<td>0.7</td>
</tr>
<tr>
<td>Return to OR during admission&lt;sup&gt;**&lt;/sup&gt;</td>
<td>-11</td>
<td>Oesophagus</td>
<td>1.5</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Gastroduodenum</td>
<td>1.4</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Small bowel</td>
<td>0.9</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Large bowel</td>
<td>1.4</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Vascular</td>
<td>1.3</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Coughing&lt;sup&gt;**&lt;/sup&gt;</td>
<td>1.4</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Wound infection&lt;sup&gt;**&lt;/sup&gt;</td>
<td>1.9</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Scoring and related risk</th>
<th>Score</th>
<th>Risk of SWD</th>
<th>Score</th>
<th>Risk of SWD</th>
<th>Score</th>
<th>Risk of SWD</th>
</tr>
</thead>
<tbody>
<tr>
<td>11-14</td>
<td>5%</td>
<td>0-2</td>
<td>0.1%</td>
<td>Range 0 to 25.7</td>
<td>Higher value predicts higher risk</td>
<td></td>
</tr>
<tr>
<td>&gt;14</td>
<td>10%</td>
<td>2-4</td>
<td>0.7%</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>4-6</td>
<td>5.5%</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>6-8</td>
<td>26.2%</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>&gt;8</td>
<td>66.5%</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<sup>*</sup>Intra-operative risk factors
<sup>**</sup>Post-operative risk factors
COPD: chronic obstructive pulmonary disease; CVA: cerebrovascular accident; OR: operating room; PGY: postgraduate year
Appendix 4. Interventions for reduction of risk of surgical site complications such as SWD and SSI

<table>
<thead>
<tr>
<th>Intervention</th>
<th>Planning</th>
<th>Pre-operative</th>
<th>Intra-operative</th>
<th>Post-operative</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Education of patient/carer/family and management of expectations</td>
<td>✔️</td>
<td>✔️</td>
<td>✔️</td>
<td></td>
<td>▪ Patients should be advised on appropriate levels of activity, hygiene measures, signs and symptoms of SWD and SSI, and when and who to contact with problems</td>
</tr>
<tr>
<td>Assessment and management/optimisation of comorbidities – e.g. obesity, malnutrition, diabetes mellitus, COPD, anaemia, cardiovascular disease</td>
<td>✔️</td>
<td>✔️</td>
<td>✔️</td>
<td></td>
<td>▪ Amelioration or removal of patient-related modifiable risk factors for SWD may reduce risk of SWD</td>
</tr>
<tr>
<td>Screening for nasal carriage of <em>Staphylococcus aureus</em> and decolonisation according to local protocol – e.g. test patients undergoing cardiac surgery or surgery involving an implant (e.g. arthroplasty or breast implant) and those who are healthcare workers or institutional residents</td>
<td>✔️</td>
<td></td>
<td></td>
<td></td>
<td>▪ Nasal carriage of <em>S. aureus</em> increases the risk of SSI after major heart surgery, breast reconstruction and implant surgery</td>
</tr>
<tr>
<td>Management of bleeding/thrombotic risk in patients on oral anticoagulants</td>
<td>✔️</td>
<td>✔️</td>
<td>✔️</td>
<td></td>
<td>▪ Management will depend on the anticoagulant in use, reason for use, risk of bleeding, procedure type and urgency, but may include cessation of the anticoagulant or replacement with a shorter acting agent</td>
</tr>
<tr>
<td>Consider nutritional supplementation</td>
<td>✔️</td>
<td>✔️</td>
<td></td>
<td></td>
<td>▪ WHO guidelines on the prevention of SSI suggest consideration of the administration of oral or enteral nutritional supplementation with multiple nutrient-enhanced formulas (containing arginine, glutamine, omega-3 fatty acids and/or nucleotides) in underweight patients who undergo major surgical operations</td>
</tr>
<tr>
<td>Use of an operative safety checklist – e.g. WHO Surgical Safety Checklist</td>
<td>✔️</td>
<td>✔️</td>
<td></td>
<td></td>
<td>▪ A systematic review and meta-analysis concluded that evidence is suggestive that use of the WHO Surgical Safety Checklist reduces post-operative complications</td>
</tr>
<tr>
<td>Maintenance of normothermia (i.e. avoidance of hypothermia, unless otherwise indicated)</td>
<td>✔️</td>
<td>✔️</td>
<td></td>
<td></td>
<td>▪ Inadvertent peri-operative hypothermia impacts wound healing</td>
</tr>
<tr>
<td>Monitor and control blood glucose of patients with diabetes mellitus</td>
<td>✔️</td>
<td>✔️</td>
<td></td>
<td></td>
<td>▪ Blood glucose levels of diabetic patients should be monitored and controlled to &lt;11mmol/l or &lt;200mg/dl</td>
</tr>
<tr>
<td>Showering or bathing by patient on day of surgery using plain or antimicrobial soap/cleanser</td>
<td>✔️</td>
<td></td>
<td></td>
<td></td>
<td>▪ Good clinical practice, but effect on surgical site complication rates is unclear and ideal type of soap/cleanser is not known</td>
</tr>
<tr>
<td>Use of clippers (rather than a razor) for hair removal</td>
<td>✔️</td>
<td></td>
<td></td>
<td></td>
<td>▪ Hair should only be removed if necessary: a meta-analysis has shown that hair removal does not reduce SSI rates; however, when hair is removed, clipping significantly reduces SSI rate in comparison with shaving</td>
</tr>
<tr>
<td>Location of heparin injection sites away from operative site</td>
<td>✔️</td>
<td>✔️</td>
<td></td>
<td></td>
<td>▪ Haematoma is more common if the heparin injection site is relatively close to the incision</td>
</tr>
<tr>
<td>Management of hydration/fluid levels to produce normovolaemia, while avoiding fluid overload and hypovolaemia</td>
<td>✔️</td>
<td>✔️</td>
<td></td>
<td></td>
<td>▪ Fluid overload may cause soft tissue oedema which may impair tissue oxygenation and wound healing; hypovolaemia may cause hypoxia</td>
</tr>
<tr>
<td>Maintenance of adequate tissue perfusion</td>
<td>✔️</td>
<td>✔️</td>
<td></td>
<td></td>
<td>▪ Haemodynamic goal-directed therapy (titration of fluids and isotropic drugs to reach target cardiac output and oxygen delivery) appears to reduce SSI</td>
</tr>
<tr>
<td>Timely administration of prophylactic antibiotics as indicated by local guidelines</td>
<td>✔️</td>
<td>✔️</td>
<td></td>
<td></td>
<td>▪ Antibiotics should be administered within the optimal time (often within 120 minutes before incision) according to the pharmacokinetics of the antibiotics in use</td>
</tr>
<tr>
<td>Administration of antifibrinolytic agents as indicated by local guidelines</td>
<td>✔️</td>
<td>✔️</td>
<td></td>
<td></td>
<td>▪ Antifibrinolytic agents, e.g. tranexamic acid and aprotinin, have been found to significantly reduce the need for blood transfusion</td>
</tr>
</tbody>
</table>
## Appendix 4. Continued

<table>
<thead>
<tr>
<th>Intervention</th>
<th>✔</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Compliance with hygiene measures by operating room personnel</td>
<td>✔</td>
<td>- For example, removal of hand jewellery, artificial nails and nail polish, covering hair, face masks, operating room suits, surgical hand/forearm preparation, sterile gloves and gowns10</td>
</tr>
<tr>
<td>Minimisation of operating room traffic</td>
<td>✔</td>
<td>- SSI usually originates from the patient’s own flora, but airborne microbes (the level of which is directly proportional to the number of people in the operating room) may play a role214</td>
</tr>
<tr>
<td>Optimal oxygenation</td>
<td>✔</td>
<td>- Supplemental oxygen reduces SSI occurrence215 - Patients should receive oxygen intra-operatively and, ideally, for 2–6 hours post-operatively142</td>
</tr>
<tr>
<td>Skin preparation with an antiseptic immediately prior to incision</td>
<td>✔</td>
<td>- It is not clear which is the most effective antiseptic solution for skin preparation215. However, the WHO recommends alcohol-based antiseptic solutions based on chlorhexidine gluconate (CHG)214</td>
</tr>
<tr>
<td>If an incise drape is necessary, use an iodophor-impregnated drape</td>
<td>✔</td>
<td>- Iodophor-impregnated incise drapes should not be used on patients who are allergic to iodine125 - In a comparison with standard incise drapes, patients who received iodophor-impregnated incise drapes had a significantly lower SSI rate106</td>
</tr>
<tr>
<td>Use of excellent surgical technique with gentle handling of tissues, meticulous control of bleeding and avoidance of dead space</td>
<td>✔</td>
<td>- Tissue trauma, poor haemostasis and failure to obliterate dead space may increase risk of SSI and impede wound healing215</td>
</tr>
<tr>
<td>Avoidance of tension across incision</td>
<td>✔</td>
<td>- High incisional tension increases the risk of SWD25</td>
</tr>
<tr>
<td>Use of wound edge protectors/guards during laparotomy</td>
<td>✔</td>
<td>- Wound protectors decrease the incidence of SSI in abdominal surgery20,172</td>
</tr>
<tr>
<td>Intra-operative wound irrigation</td>
<td>✔</td>
<td>- Intra-operative wound irrigation reduces SSI rates, with the most marked effect in colorectal surgery208 - WHO and CDC guidelines recommend the use of aqueous iodophor solution124,173</td>
</tr>
<tr>
<td>Change of gloves during procedure and/or before closure of wound, double gloving</td>
<td>✔</td>
<td>- Widely practiced, especially for high risk/contaminated procedures, but effect on SSI rates is unclear215</td>
</tr>
<tr>
<td>Senior/experienced surgeon performing closure</td>
<td>✔</td>
<td>- There is an inverse association between the level of experience of a surgeon and SSI rates: autonomously performed closure has been found to have a significantly lower SSI rate than closure performed under supervision199 - In comparison with a more experienced surgeon, surgery performed by a postgraduate year 4 surgeon is associated with an increased rate of SWD28</td>
</tr>
<tr>
<td>Use of gentamicin-impregnated collagen sponges</td>
<td>✔</td>
<td>- Reduce rates of SSI in cardiac, colorectal and femoropopliteal bypass surgery209–211</td>
</tr>
<tr>
<td>Use of triclosan-coated sutures</td>
<td>✔</td>
<td>- Triclosan-coated sutures should be considered because they reduce rates of SSI203,212</td>
</tr>
<tr>
<td>Covering of the incision(s) with a dry absorbent sterile dressing under sterile conditions and before the patient leaves the operating room</td>
<td>✔</td>
<td>- Dressings provide a physical barrier to external contamination216</td>
</tr>
<tr>
<td>Consider prophylactic NPWT (e.g. single-use NPWT) for patients at increased risk of SSI or SWD</td>
<td>✔</td>
<td>- The WHO recommends the prophylactic use of NPWT on closed surgical incisions in high-risk patients to prevent SSI143</td>
</tr>
<tr>
<td>Maintenance of the dressing over the incision for at least 48 hours unless there are signs and symptoms indicating earlier inspection is warranted</td>
<td>✔</td>
<td>- Epithelialisation of surgical wounds is usually complete, i.e. the wound is sealed, within 48 hours. Therefore, dressings should be inspected regularly but left in place for at least first 48 hours post-operatively143 - If a dressing change is required before 48 hours, the dressing should be changed using an aseptic technique215</td>
</tr>
<tr>
<td>Cryotherapy (e.g. application of ice) and compression</td>
<td>✔</td>
<td>- Cryotherapy and compression aim to aid healing by reducing oedema that may be impairing tissue perfusion216,272 - A retrospective study of patients who underwent total ankle arthroplasty reported that a compression wrap protocol reduced wound-related complications (a composite endpoint that included SWD)291 - Cryotherapy is widely used for pain relief following orthopaedic surgery and may be combined with compression therapy216</td>
</tr>
<tr>
<td>Visitor restrictions and hygiene measures – e.g. hand cleansing/protective clothing as appropriate</td>
<td>✔</td>
<td>- An SSI bundle for patients undergoing cardiac surgery that included visitor restrictions resulted in a lower incidence of SSI275</td>
</tr>
<tr>
<td>Monitor incision for healing progress and signs/symptoms of dehiscence or infection</td>
<td>✔</td>
<td>- Early recognition of problems followed by appropriate interventions is likely to improve longer term outcomes</td>
</tr>
<tr>
<td>Patient Reported Outcome/Experience Measures (PROMS/PREMS) or questionnaires</td>
<td>✔</td>
<td>- Increasingly used for monitoring and may be linked to reimbursement in some healthcare systems205</td>
</tr>
<tr>
<td>Perform surveillance of post-operative wound complications and compliance with surgical wound complication reduction bundles</td>
<td>✔</td>
<td>- Active surveillance may decrease SSI rates276 - Aids feedback to individual surgeons and team members and monitoring of trends/effect of implementation of measures to reduce SWD/SSI207</td>
</tr>
</tbody>
</table>
Appendix 4. Continued

Post-operative complications and compliance with surgical

Perform surveillance of post-operative wound measures – e.g. hand cleansing/

Visitor restrictions and hygiene

compression

there are signs and symptoms indicating

Use NPWT) for patients at increased risk

closure

Senior/ experienced surgeon performing

before closure of wound; double gloving

Change of gloves during procedure and/ or

✔

Use of wound edge protectors/ guards

✔ ✔

Intra-operative wound irrigation reduces SSI rates, with the most marked effect in colorectal

Iodophor-impregnated incise drapes should not be used on patients who are allergic to iodine121

It is not clear which is the most effective antiseptic solution for skin preparation173. However, the

For example, removal of hand jewellery, artificial nails and nail polish, covering hair, face

Aids feedback to individual surgeons and team members and monitoring of trends/ effect of

SUPPLEMENT

Tissue trauma, poor haemostasis and failure to obliterate dead space may increase risk of

An SSI bundle for patients undergoing cardiac surgery that included visitor restrictions

4 surgeon is associated with an increased rate of SWD79

In comparison with a more experienced surgeon, surgery performed by a postgraduate year

there is an inverse association between the level of experience of a surgeon and SSI rates:

Widely practiced, especially for high risk/ contaminated procedures, but effect on SSI rates

Ensure wounds are not overly tensioned

For example, removal of hand jewellery, artificial nails and nail polish, covering hair, face

✔

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