# Surgery on Wrong Body Part

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<tr>
<th>Event Category</th>
<th>Full Term</th>
<th>Short Term</th>
<th>Specifications</th>
<th>Standard</th>
</tr>
</thead>
<tbody>
<tr>
<td>Surgical</td>
<td>Surgery or other invasive procedure performed on the wrong site</td>
<td>Surgery on wrong body part</td>
<td>Defined as any surgery or other invasive procedure performed on a body part or site that is not consistent with the correctly documented informed consent for that patient. Surgery or other invasive procedure includes, but is not limited to, endoscopies, lens implants, lesion removal, injection into joints. Excludes emergent situations that occur in the course of surgery or other invasive procedure and/or whose exigency precludes obtaining informed consent.</td>
<td>NQF 1A</td>
</tr>
</tbody>
</table>

**Implementation Guidance**

Wrong site surgery or invasive procedure, corrected during the procedure, is still a wrong site procedure if the surgery/procedure had begun, based on the definition in glossary.

This event is intended to capture instances of:

- surgery or other invasive procedure on the right body part but on the wrong location/site on the body; example, left/right (appendages and/or organs), wrong digit, level (spine), stent placed in wrong iliac artery, steroid injection into wrong knee, biopsy of wrong mole, burr hole on wrong side of skull;
- delivery of fluoroscopy or radiotherapy to the wrong region of the body;
- use of incorrectly placed vascular catheters;
- use of incorrectly placed tubes (for example, feeding tubes placed in the lung or ventilation tubes passed into the esophagus);
This event is not intended to capture:

- changes in plan upon entry into the patient with discovery of pathology in close proximity to the intended place where risk of a second surgery or procedure outweighs the benefit of patient consultation or unusual physical configuration (e.g., adhesions, spine level/extra vertebrae).

**examples**

*actual sentinel event:* A hand surgeon performs trigger finger surgery on the wrong finger. Before applying the dressing, the surgeon realizes the mistake. He then performs the procedure on the correct finger.
surgery on wrong patient

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<tbody>
<tr>
<td>surgical</td>
<td>surgery or other invasive procedure performed on the wrong patient</td>
<td>surgery on wrong patient</td>
<td>Defined as any surgery or invasive procedure on a patient that is not consistent with the correctly documented informed consent for that patient. Surgery or other invasive procedure includes, but is not limited to, endoscopies, lens implants, lesion removal, injection into joints.</td>
<td>NQF 1B</td>
</tr>
</tbody>
</table>

**implementation guidance**

This event is intended to capture:

- surgical procedures (whether or not completed) initiated on one patient intended for a different patient.

**examples**

*risk thereof sentinel event: A 42 year-old male is admitted for pneumonia. The patient is not planning to have any surgical procedure. The patient’s roommate is scheduled to have an elective cyst removal. A nurse confuses the bed numbers and takes the patient with the pneumonia to the operating room. The patient is prepped and placed under general anesthesia. Following the administration of anesthesia, the mistake is identified. There has been no skin perforation, yet there was a risk of surgery on the wrong patient.*
### Wrong Surgical Procedure

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</tr>
</thead>
<tbody>
<tr>
<td>Surgical</td>
<td>Wrong surgical or other invasive procedure performed on a patient</td>
<td>Wrong surgical procedure</td>
<td>Defined as any surgical or other invasive procedure performed on a patient that is not consistent with the correctly documented informed consent for that patient. Surgery or other invasive procedure includes, but is not limited to, endoscopies, lens implants, lesion removal, injection into joints. Excludes emergent situations that occur in the course of surgery or other invasive procedures and/or whose exigency precludes obtaining informed consent.</td>
<td>NQF 1C</td>
</tr>
</tbody>
</table>

### Implementation Guidance

This event is intended to capture:

- Any surgical procedure performed incorrectly.

This event is not intended to capture:

- Changes in plan upon surgical entry into the patient with discovery of pathology in close proximity to the intended place where risk of a second surgery/procedure outweighs benefit of patient consultation, or unusual physical configuration (for example adhesions, spine level/extra vertebrae).

This event is not intended to capture; changes in plan upon entry into the patient with discovery of pathology in close proximity to the intended place where risk of a second surgery/procedure outweighs benefit of patient consultation, or unusual physical configuration (for example adhesions, spine level/extra vertebrae).

### Examples
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<tbody>
<tr>
<td>surgical</td>
<td>unintended retention of a foreign object in a patient after surgery or other invasive procedure</td>
<td>retained foreign object</td>
<td>Excludes a) objects present prior to surgery or other invasive procedure that are intentionally left in place; b) objects intentionally implanted as part of a planned intervention; and c) objects not present prior to surgery/procedure that are intentionally left in when the risk of removal exceeds the risk of retention (such as microneedles, broken screws).</td>
<td>NQF 1D</td>
</tr>
</tbody>
</table>

**implementation guidance**

This event is intended to capture:

- occurrences of unintended retention of objects at any point after the surgery/procedure ends regardless of setting (post anesthesia recovery unit, surgical suite, emergency department, patient bedside) and regardless of whether the object is to be removed after discovery.
- unintentionally retained objects (including such things as wound material, sponges, catheter tips, trocars, guide wires) in all applicable settings.

**examples**

*risk thereof sentinel event*: A 29 year-old female presents in active labor. The patient is taken to labor and delivery. During labor, the baby is found to be in fetal distress and a leg is identified at the cervical os. The mother is taken to the operating room emergently. A C-section is performed and a healthy baby is delivered. During the initial count, a missing lap is identified. The cavity is examined, and the count is repeated, but the missing lap remains. Radiologic evaluation does not reveal the lap either. The incision is closed. While the patient is transferred to the recovery room, an end of a lap is noted in the vaginal canal and is successfully removed.
### intra- or post-operative death

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</thead>
<tbody>
<tr>
<td>surgical</td>
<td>intraoperative or immediately postoperative/postprocedure death in an ASA Class I patient</td>
<td>intra- or post-operative death</td>
<td>Includes all ASA Class I patient deaths in situations where anesthesia was administered; the planned surgical procedure may or may not have been carried out.</td>
<td>NQF 1E</td>
</tr>
<tr>
<td></td>
<td>ASA 1: No organic pathology or patients in whom the pathological process is localized and does not cause any systemic disturbance or abnormality</td>
<td></td>
<td>Immediately post-operative means within 24 hours after surgery or other invasive procedure was completed or after administration of anesthesia (if surgery/procedure was not completed).</td>
<td></td>
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<tr>
<td></td>
<td>(Unexpected death in other ASA Class patients would be captured in OTHER category)</td>
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<td></td>
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</tbody>
</table>

### implementation guidance

This event is intended to capture:

- ASA Class I patient death associated with the administration of anesthesia whether or not the planned surgical procedure was carried out.

### examples

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<tr>
<th>examples</th>
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</table>
contaminated drug, device, or biologic

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<tbody>
<tr>
<td>product or device</td>
<td>patient death or serious injury associated with the use of contaminated drugs, devices, or biologics provided by the healthcare setting</td>
<td>contaminated drug, device, or biologic</td>
<td>Includes contaminants in drugs, devices, or biologics regardless of the source of contamination and/or product. Includes threat of disease that changes patient’s risk status for life requiring medical monitoring not needed before the event</td>
<td>NQF 2A</td>
</tr>
</tbody>
</table>

**Implementation guidance**

This event is intended to capture:

- contaminations that can be seen the naked eye or with use of detection mechanisms in general use. These contaminations are to be reported at such time as they become known to the provider or healthcare organization. Contaminants may be physical, chemical, or biological in nature. Not all contaminations can be seen with the naked eye (e.g., acid testing, mass spectrometry, and tests that signal changes in pH or glucose levels). Contamination that is inferred and changes risk status for life (e.g., consider a syringe or needle contaminated once it has been used to administer medication to a patient by injection or via connection to a patient’s intravenous infusion bag or administration set).

This event is intended to capture:

- administration of contaminated vaccine or medication (e.g., intramuscular antibiotic);
- serious infection from contaminated drug or device used in surgery or an invasive procedure (e.g., a scalpel);
- occurrences related to use of improperly cleaned or maintained device.

**Examples**

Hepatitis C is an example of a disease that changes a patient’s risk status for life, requiring ongoing treatment.
### Device Failure

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<tr>
<td>Product or Device</td>
<td>Patient death or serious injury associated with the use or function of a device in patient care, in which the device is used or functions other than as intended</td>
<td>Device failure</td>
<td>Includes, but is not limited to, catheters, drains and other specialized tubes, infusion pumps, ventilators, and procedural and monitoring equipment.</td>
<td>NQF 2B</td>
</tr>
</tbody>
</table>

**Implementation Guidance**

This event is intended to capture:

- occurrences whether or not the use is intended or described by the device manufacturers’ literature

**Examples**
### air embolism

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<tbody>
<tr>
<td>product or device</td>
<td>patient death or serious injury associated with intravascular air embolism that occurs while being cared for in a healthcare setting</td>
<td>air embolism</td>
<td>Excludes death or serious injury associated with neurosurgical procedures known to present a high risk of intravascular air embolism.</td>
<td>NQF 2C</td>
</tr>
</tbody>
</table>

**implementation guidance**

This event is intended to capture:

- high-risk procedures, other than neurosurgical procedures, that include, but are not limited to, procedures involving the head and neck, vaginal delivery and cesarean section, spinal instrumentation procedures, and liver transplantation;
- low-risk procedures, including those related to lines placed for infusion of fluids in vascular space.

**examples**

*risk thereof sentinel event:* Air embolism includes procedures that have a small but known risk—again confirming the fact that risk does not imply expected outcome.
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<tbody>
<tr>
<td>patient protection</td>
<td>discharge or release of a patient/resident of any age, who is unable to make decisions, to other than an authorized person</td>
<td>discharge to wrong person</td>
<td></td>
<td>NQF 3A</td>
</tr>
</tbody>
</table>

**implementation guidance**

The terms “authorized” and “decision-making capacity” are defined in the glossary. Release to “other than an authorized person” includes removing the patient/resident without specific notification and approval by staff, even when the person is otherwise authorized.

Examples of individuals who do not have decision-making capacity include: newborns, minors, adults with Alzheimer’s.

Individual healthcare organizations or other relevant jurisdictional authorities may have specific requirements for assessing decisionmaking capacity.

**examples**

*risk thereof sentinel event*: Mrs. M delivers a healthy baby boy. In preparation for discharge home, Mrs. M signs all documents and is ready to go. At the time of departure, Mrs. G’s baby is delivered to Mrs. M. Upon arrival of the baby, Mrs. M states that the baby is not hers. Nursing is notified, the error is acknowledged, and the correct baby goes home with Mrs. M. In this situation, multiple corrective measures failed to recognize the wrong baby.
# elopement

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<tbody>
<tr>
<td>patient protection</td>
<td>patient death or serious injury associated with patient elopement (disappearance)</td>
<td>elopement</td>
<td>Includes events that occur after the individual presents him/herself for care in a healthcare setting. Excludes events involving competent adults with decision-making capacity who leave against medical advice or voluntarily leave without being seen.</td>
<td>NQF 3B</td>
</tr>
</tbody>
</table>

## Implementation guidance

The term “elopement” and “competent” adult should be interpreted in accordance with prevailing legal standards in applicable jurisdictions.

This event is not intended to capture:

- death or serious injury that occurs (after the patient is located) due to circumstances unrelated to the elopement.

## Examples
### Event Category: Suicide

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</tr>
</thead>
<tbody>
<tr>
<td>Patient Protection</td>
<td>Patient suicide, attempted suicide, or self-harm that results in serious</td>
<td>Suicide</td>
<td>Includes events that result from patient actions after they present themselves for care in a healthcare setting.</td>
<td>NQF 3C</td>
</tr>
<tr>
<td></td>
<td>injury, while being cared for in a healthcare setting</td>
<td></td>
<td>Excludes deaths resulting from self-inflicted injuries that were the reason for admission/presentation to the healthcare facility.</td>
<td></td>
</tr>
</tbody>
</table>

### Implementation Guidance

This event is not intended to capture patient suicide or attempted suicide when the patient is not physically present in the “healthcare setting” as defined in the glossary.

### Examples

<table>
<thead>
<tr>
<th>Example 1</th>
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<table>
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<tr>
<th>Example 2</th>
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</table>
### Medication Error

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</tr>
</thead>
<tbody>
<tr>
<td>Care Management</td>
<td>Patient death or serious injury associated with a medication error (e.g., errors involving the wrong drug, wrong dose, wrong patient, wrong time, wrong rate, wrong preparation, or wrong route of administration)</td>
<td>Medication error</td>
<td>Excludes reasonable differences in clinical judgment on drug selection and dose. Includes, but is not limited to, death or serious injury associated with: a) over- or under-dosing; b) administration of a medication to which a patient has a known allergy or serious contraindication, c) drug-drug interactions for which there is known potential for death or serious disability, and improper use of single-dose/single-use and multi-dose medication vials and containers leading to death or serious injury as a result of dose adjustment problems.</td>
</tr>
</tbody>
</table>

### Implementation Guidance

This event is intended to capture:

- the most serious medication errors including occurrences in which a patient receives a medication for which there is a contraindication, or a patient known to have serious allergies to specific medications/agents, receives those medications/agents, resulting in serious injury or death. These events may occur as a result of failure to collect information about contraindications or allergies, failure to review such information available in information systems, failure of the organization to ensure availability of such information and prominently display such information within information systems, or other system failures that are determined through investigation to be the cause of the adverse event;
- occurrences in which a patient dies or suffers serious injury as a result of failure to administer a prescribed medication;
- occurrences in which a patient is administered an over- or under-dose of a medication including insulin, heparin, and any other high alert medication including but not limited to medications listed on the Institute for Safe Medication Practices “High Alert Medication List”;
- occurrences in which a patient dies or suffers serious injury as a result of the wrong administration technique.

This event is not intended to capture:

- patient death or serious injury associated with allergies that could not reasonably have been known or discerned in advance of the event. These unexpected deaths would be captured in “other” category.
**examples**

*actual sentinel event*: A terminally ill cancer patient receives a 10-fold overdose of morphine and dies within 24 hours. The patient was lucid prior to the overdose; after the overdose, the patient is comatose and never recovers consciousness before dying. Despite the patient’s terminal condition this is a medication error event if the patient was resuscitated and lived, this would still be a sentinel event.
transfusion error

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</thead>
<tbody>
<tr>
<td>care management</td>
<td>patient death or serious injury associated with unsafe administration of blood products</td>
<td>transfusion error</td>
<td></td>
<td>NQF 4B</td>
</tr>
</tbody>
</table>

**Implementation guidance**

Unsafe administration includes, but is not limited to, hemolytic reactions and administering: a) blood or blood products to the wrong patient; b) the wrong type; or c) blood or blood products that have been improperly stored or handled.

This event is not intended to capture:

- patient death or serious injury associated with organ rejection other than those attributable to a hyperacute hemolytic reaction
- patient death or injury when cause is not detectable by ABO/HLA matching.
- These deaths would be categorized in ‘other’ category.

**Examples**


maternal labor or delivery

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<tbody>
<tr>
<td>care management</td>
<td>maternal death or serious injury associated with labor or delivery in a pregnancy while being cared for in a healthcare setting</td>
<td>maternal labor or delivery</td>
<td></td>
<td>NQF 4C</td>
</tr>
</tbody>
</table>

**implementation guidance**

This event is not intended to create a new obligation. The organization’s obligation, under this event, is to report only maternal death or serious injury associated with labor or delivery in a pregnancy when made aware of the maternal death or serious injury either by readmittance or by the patient’s family.

**examples**


### neonate labor or delivery

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<tbody>
<tr>
<td>care management</td>
<td>death or serious injury of neonate associated with labor or delivery in a pregnancy</td>
<td>neonate labor or delivery</td>
<td>Includes, for the office-based surgery, birthing center or “home” setting, unplanned admission to an inpatient setting within 24 hours of delivery</td>
<td>NQF 4D</td>
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### implementation guidance

### examples
**fall**

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<tr>
<td>care management</td>
<td>patient death or serious injury associated with a fall while being cared for in a healthcare setting</td>
<td>fall</td>
<td>Includes but is not limited to fractures, head injuries, and intracranial hemorrhage</td>
<td>NQF 4E</td>
</tr>
</tbody>
</table>

**implementation guidance**

Of note, an assessment that identifies patients at “risk” of fall, findings of risk accompanied by organizationally defined measures to be taken when risk is identified could be useful in both prevention and event analysis.

This category is not intended to include slips and trips.

**examples**

*risk thereof sentinel event*: A 44-year-old morbidly obese male is admitted for a knee replacement. At the time of diagnosis, patient is assessed as a high risk for a fall and appropriate measures are recommended. The measures include 2-person assist, call light, close proximity to nursing, and room signage. During hospitalization, the patient uses the call light to alert nursing that he needs to use the restroom. Only 1 nurse responds and assists the patient to the restroom. The nurse then leaves the patient on the toilet to get the second nurse to assist. The patient falls from the toilet and strikes his hip and leg. Two nurses then return to the room and assist the patient back to bed. The patient complains of sore hip and X-rays are done, but no fracture is identified. This is a process failure and falls into the risk thereof category and thus needs to be reported. A fall does not need to result in permanent loss of function or death as a direct result of injuries sustained by the fall to qualify as a sentinel event.

*actual sentinel event*: A patient is admitted with a diagnosis of lithium intoxication. The patient falls while in the hospital resulting in a fracture of the right humerus. Post X-ray, the patient is placed in a sling, and no other intervention is required. Hospital reasons that this is not a sentinel event because the injury sustained as a result of the fall did not lead to major permanent loss of function or death. The sentinel event definition contains no such criterion by which an event must result in permanent loss or death to qualify as a sentinel event. Permanent loss of function or death are not required outcomes for an event to be considered a sentinel event.
## Pressure Ulcer

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<tbody>
<tr>
<td>Care Management</td>
<td>Any stage 3, stage 4, and unstageable pressure ulcers acquired after admission/presentation to a healthcare setting</td>
<td>Pressure Ulcer</td>
<td>Excludes progression from Stage 2 to Stage 3 if Stage 2 was recognized upon admission and excludes pressure ulcers that develop in areas where deep tissue injury is documented as present on admission/presentation.</td>
<td>NQF 4F</td>
</tr>
</tbody>
</table>

### Implementation Guidance

### Examples

*Actual sentinel event*: A 68 year-old male with history of acute and chronic respiratory failure and diagnosis of lung cancer presents. Skin assessment on admission is clear. Nursing notes revealed redness on hospital day 14. On hospital day 23, patient found to have stage 4 pressure ulcer. Despite patient’s diagnosis and co-morbidities, a pressure ulcer is not expected.
### wrong sperm or egg

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</thead>
<tbody>
<tr>
<td>care management</td>
<td>artificial insemination with the wrong donor sperm or wrong egg</td>
<td>wrong sperm or egg</td>
<td></td>
<td>NQF 4G</td>
</tr>
</tbody>
</table>

### implementation guidance

The organization’s obligation is to report the event when it is made aware of the occurrence.

### examples

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<table>
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### lost specimen

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<tbody>
<tr>
<td>care management</td>
<td>patient death or serious injury resulting from the irretrievable loss of an irreplaceable biological specimen</td>
<td>lost specimen</td>
<td>Includes events where specimens are misidentified, where another procedure cannot be done to produce a specimen</td>
<td>NQF 4H</td>
</tr>
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<td></td>
<td>Includes progression of an undiagnosed disease or threat of disease that changes the patient’s risk status for life, requiring monitoring not needed before the event</td>
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</tbody>
</table>

#### implementation guidance
This event is not intended to capture:

- procedures where the specimen was properly handled, but the specimen proved to be nondiagnostic.

#### examples

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**electric shock**

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<tbody>
<tr>
<td>environment</td>
<td>patient or staff death or serious injury associated with an electric shock in the course of a patient care process in a healthcare setting</td>
<td>electric shock</td>
<td>Excludes events involving patients during planned treatments such as electric countershock/elective cardioversion.</td>
<td>NQF 5A</td>
</tr>
</tbody>
</table>

**implementation guidance**

This event is intended to capture:

- patient death or injury associated with unintended electric shock during the course of care or treatment;
- staff death or injury associated with unintended electric shock while carrying out duties directly associated with a patient care process, including preparing for care delivery.

This event is not intended to capture:

- patient death or injury associated with emergency defibrillation in ventricular fibrillation or with electroconvulsine therapies;
- injury to staff who are not involved in patient care.

**examples**
### wrong or contaminated gas

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<tbody>
<tr>
<td>environment</td>
<td>any incident in which systems designated for oxygen or other gas to be delivered to a patient contains no gas, the wrong gas, or are contaminated by toxic substances</td>
<td>wrong or contaminated gas</td>
<td></td>
<td>NQF 5B</td>
</tr>
</tbody>
</table>

**implementation guidance**

This event is intended to capture:

- events in which the line is attached to a reservoir distant from the patient care unit or in a tank near the patient such as E-cylinders, anesthesia machines.

**examples**


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<tbody>
<tr>
<td>environment</td>
<td>patient or staff death or serious injury associated with a burn incurred from any source in the course of a patient care process in a healthcare setting</td>
<td>burn</td>
<td></td>
<td>NQF 5C</td>
</tr>
</tbody>
</table>

**implementation guidance**

This event is intended to capture burns that result from:

- operating room flash fires, including second-degree burn in these cases;
- hot water;
- sunburn in the patient with decreased ability to sense pain;
- smoking in the patient care environment.

**examples**
**restraint**

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<tbody>
<tr>
<td>environment</td>
<td>patient death or serious injury associated with the use of physical restraints or bedrails while being cared for in a healthcare setting</td>
<td>restraint</td>
<td></td>
<td>NQF 5D</td>
</tr>
</tbody>
</table>

**implementation guidance**

This event is intended to capture:

- instances where physical restraints are implicated in the death, e.g., lead to strangulation/entrapment, etc.

**examples**


<table>
<thead>
<tr>
<th>event category</th>
<th>full term</th>
<th>short term</th>
<th>specifications</th>
<th>standard</th>
</tr>
</thead>
<tbody>
<tr>
<td>radiologic</td>
<td>death or serious injury of a patient or staff associated with the introduction of a metallic object into the MRI area</td>
<td>introduction of metallic object into MRI area</td>
<td>Includes events related to material inside the patient’s body or projectiles outside the patient’s body.</td>
<td>NQF 6A</td>
</tr>
</tbody>
</table>

**implementation guidance**

This event is intended to capture:

- retained foreign objects
- external projectiles
- pacemakers

**examples**
impersonation of healthcare provider

<table>
<thead>
<tr>
<th>event category</th>
<th>full term</th>
<th>short term</th>
<th>specifications</th>
<th>standard</th>
</tr>
</thead>
<tbody>
<tr>
<td>criminal</td>
<td>any instance of care ordered by or provided by someone impersonating a physician, nurse, pharmacist, or other licensed healthcare provider</td>
<td>impersonation of healthcare provider</td>
<td></td>
<td>NQF 7A</td>
</tr>
</tbody>
</table>

**implementation guidance**

This event is intended to capture:

- those without licensure to provide the care given;
- those with licensure who represent themselves and act beyond the scope of their licensure.

It is not intended to capture individuals who are practicing within the scope of their license on whom patients or others mistakenly bestow titles beyond that scope when such is not encouraged by the provider.

**examples**
abduction

<table>
<thead>
<tr>
<th>event category</th>
<th>full term</th>
<th>short term</th>
<th>specifications</th>
<th>standard</th>
</tr>
</thead>
<tbody>
<tr>
<td>criminal</td>
<td>abduction of a patient/resident of any age</td>
<td>abduction</td>
<td></td>
<td>NQF 7B</td>
</tr>
</tbody>
</table>

**implementation guidance**

This event is intended to capture:

- removal of a patient/resident, who does not have decisionmaking capacity, without specific notification and approval by staff even when the person is otherwise authorized to be away from the setting.

Examples of individuals who do not have decisionmaking capacity include: newborns, minors, adults with Alzheimer’s.

**examples**

---

2012-01-01
### Sexual Assault

<table>
<thead>
<tr>
<th>Event Category</th>
<th>Full Term</th>
<th>Short Term</th>
<th>Specifications</th>
<th>Standard</th>
</tr>
</thead>
<tbody>
<tr>
<td>Criminal</td>
<td>Sexual abuse/assault on a patient or staff member within or on the grounds of a healthcare setting</td>
<td>Sexual assault</td>
<td></td>
<td>NQF 7C</td>
</tr>
</tbody>
</table>

### Implementation Guidance

Language and definitions may vary based on state statute; however, the principle and intent remain regardless of language required based on jurisdiction.

### Examples
physical assault

<table>
<thead>
<tr>
<th>event category</th>
<th>full term</th>
<th>short term</th>
<th>specifications</th>
<th>standard</th>
</tr>
</thead>
<tbody>
<tr>
<td>criminal</td>
<td>death or serious injury of a patient or staff member resulting from a physical assault (i.e., battery) that occurs within or on the grounds of a healthcare setting</td>
<td>physical assault</td>
<td></td>
<td>NQF 7D</td>
</tr>
</tbody>
</table>

implementation guidance

Language and definitions may vary based on state statute (e.g., many states have existing statutes that use the terms “first degree assault” or “second degree assault” or “battery”).

examples


### CLABSI

<table>
<thead>
<tr>
<th>event category</th>
<th>full term</th>
<th>short term</th>
<th>specifications</th>
<th>standard</th>
</tr>
</thead>
<tbody>
<tr>
<td>healthcare-associated</td>
<td>primary bloodstream infection that is central line-associated</td>
<td>CLABSI</td>
<td></td>
<td></td>
</tr>
<tr>
<td>infection</td>
<td></td>
<td></td>
<td></td>
<td>AHRQ, NHSN</td>
</tr>
</tbody>
</table>

**Implementation guidance**

A simplified tool based on NHSN guidance and a CLABSI checklist are included on the next few pages.

**Examples**

*actual sentinel event*: A catheter tip isolate is *S aureus*. The blood culture isolate is coagulase-negative *staphylococci* (CNS). The hospital reasons that when CNS is found in a blood culture, it is usually a skin contaminant and the specimen is probably just contaminated and, therefore, unreliable. The case could not rule out infection from another source causing the blood-stream infection. Despite this reasoning, if a common commensal is isolated, it should not be automatically assumed that it does not meet the criteria for a CLABSI. NHSN criteria must be reviewed to help make the appropriate determination.

*actual sentinel event*: One of two cultures test positive. The hospital reasons that the positive test is probably just due to contamination since only one of two cultures test positive; therefore, the CLABSI is not proven. Despite this reasoning, it should not be assumed that just because only one of two cultures test positive that the NHSN criteria for a CLABSI are not met. NHSN criteria must be reviewed to help make the appropriate determination.
Central Line-Associated Bloodstream Infection (CLABSI) Event

**Introduction:** An estimated 248,000 bloodstream infections occur in U.S. hospitals each year\(^1\) and a large proportion of these are associated with the presence of a central vascular catheter. For the purposes of NHSN, such infections are termed central line-associated bloodstream infections (CLABSI). Bloodstream infections are usually serious infections typically causing a prolongation of hospital stay and increased cost and risk of mortality.

CLABSI can be prevented through proper management of the central line. These techniques are addressed in the CDC’s Healthcare Infection Control Practices Advisory Committee (CDC/HIPAC) *Guidelines for the Prevention of Intravascular Catheter-Related Infections, 2011.*\(^2\)

**Settings:** Surveillance will occur in any inpatient location where denominator data can be collected, which may include critical/intensive care units (ICU), specialty care areas (SCA), neonatal units including neonatal intensive care units (NICUs), stepdown units, wards, and long term care units. A complete listing of inpatient locations can be found in Chapter 15.

NOTE: Surveillance for CLABSIIs after the patient is discharged from the facility is not required, however, if discovered, these infections should be reported to NHSN. No additional central line days are reported.

**Requirements:** Surveillance for CLABSI in at least one inpatient location in the healthcare institution for at least one calendar month as indicated in the *Patient Safety Monthly Reporting Plan* (CDC 57.106).

**Definitions:** As for all infections reported to NHSN, infections associated with complications or extensions of infections already present on admission, unless a change in pathogen or symptoms strongly suggests the acquisition of a new infection, are not considered healthcare associated. Therefore, infections that become apparent within the first few days of admission must be carefully reviewed to determine whether they should be considered healthcare associated.

Primary bloodstream infections (BSI) are laboratory-confirmed bloodstream infections (LCBI) that are not secondary to an HAI meeting CDC/NHSN criteria at another body site (see criteria in Chapter 17 or a community-associated infection.) Report BSIs that are central line associated (i.e., a central line or umbilical catheter was in place at the time of, or within 48 hours before, onset of the event).

NOTE: There is no minimum period of time that the central line must be in place in order for the BSI to be considered central line associated.
Location of attribution: The inpatient location where the patient was assigned on the date of the BSI event, which is further defined as the date when the first clinical evidence appeared or the date the specimen used to meet the BSI criteria was collected, whichever came first.

EXAMPLE: Patient who had no clinical signs or symptoms of sepsis upon arrival to the Emergency Department, has a central line inserted there before being admitted to the MICU has a central line inserted in the Emergency Department and then is admitted to the MICU. Within 24 hours of admission to the MICU, patient meets criteria for BSI. This is reported to NHSN as a CLABSI for the MICU, because the Emergency Department is not an inpatient location and no denominator data are collected there.

TRANSFER RULE EXCEPTION: If a CLABSI develops within 48 hours of transfer from one inpatient location to another in the same facility, or a new facility, the infection is attributed to the transferring location. This is called the Transfer Rule and examples are shown below:

- Patient with a central line in place in the SICU is transferred to the surgical ward. Thirty six (36) hours later, the patient meets the criteria for BSI. This is reported to NHSN as a CLABSI for the SICU.
- Patient is transferred to the medical ward from the MSICU after having the central line removed. Within 24 hours, patient meets criteria for a BSI. This is reported to NHSN as a CLABSI for the MSICU.
- Patient with a central line in place is transferred from the medical ward to the coronary care ICU (CCU). After 4 days in the CCU, the patient meets the criteria for a BSI. This is reported to NHSN as a CLABSI for the CCU.
- Patient on the urology ward of Hospital A had the central line removed and is discharged home a few hours later. The IP from Hospital B calls the next day to report that this patient has been admitted to Hospital B with a BSI. This CLABSI should be reported to NHSN for, and by, Hospital A and attributed to the urology ward. No additional catheter days are reported.

Central line: An intravascular catheter that terminates at or close to the heart or in one of the great vessels which is used for infusion, withdrawal of blood, or hemodynamic monitoring. The following are considered great vessels for the purpose of reporting central-line BSI and counting central-line days in the NHSN system: Aorta, pulmonary artery, superior vena cava, inferior vena cava, brachiocephalic veins, internal jugular veins, subclavian veins, external iliac veins, common iliac veins, femoral veins, and in neonates, the umbilical artery/vein.

NOTES:

1. Neither the insertion site nor the type of device may be used to determine if a line qualifies as a central line. The device must terminate in one of these vessels or in or near the heart to qualify as a central line.

2. An introducer is considered an intravascular catheter, and depending on the location of its tip, may be a central line.
3. Pacemaker wires and other nonlumened devices inserted into central blood vessels or the heart are not considered central lines, because fluids are not infused, pushed, nor withdrawn through such devices.

4. The following devices are not considered central lines: extracorporeal membrane oxygenation (ECMO), femoral arterial catheters and Intraaortic balloon pump (IABP) devices. If you have a question about whether a device qualifies as a central line, please email us at NHSN@cdc.gov.

**Infusion:** The introduction of a solution through a blood vessel via a catheter lumen. This may include continuous infusions such as nutritional fluids or medications, or it may include intermittent infusions such as flushes or IV antimicrobial administration, or blood, in the case of transfusion or hemodialysis.

**Umbilical catheter:** A central vascular device inserted through the umbilical artery or vein in a neonate.

**Temporary central line:** A non-tunneled catheter.

**Permanent central line:** Includes
- Tunneled catheters, including certain dialysis catheters
- Implanted catheters (including ports)

**Laboratory-confirmed bloodstream infection (LCBI):** Must meet one of the following criteria:

**Criterion 1:** Patient has a recognized pathogen cultured from one or more blood cultures and organism cultured from blood is not related to an infection at another site. (See Notes 1 and 2 below.)

**Criterion 2:** Patient has at least one of the following signs or symptoms: fever (>38°C), chills, or hypotension and signs and symptoms and positive laboratory results are not related to an infection at another site and common commensal (i.e., diphtheroids [Corynebacterium spp. not C. diphtheriae], Bacillus spp. [not B. anthracis], Propionibacterium spp., coagulase-negative staphylococci [including S. epidermidis], viridans group streptococci, Aerococcus spp., Micrococcus spp.) is cultured from two or more blood cultures drawn on separate occasions.

**Criterion 3:** Patient ≤ 1 year of age has at least one of the following signs or symptoms: fever (>38°C core) hypothermia (<36°C core), apnea, or bradycardia and
signs and symptoms and positive laboratory results are not related to an infection at another site and common skin commensal (i.e., diphtheroids [Corynebacterium spp. not C. diphtheriae], Bacillus spp. [not B. anthracis], Propionibacterium spp., coagulase-negative staphylococci [including S. epidermidis], viridans group streptococci, Aerococcus spp., Micrococcus spp.) is cultured from two or more blood cultures drawn on separate occasions. (See Notes 3, 4 and 5 below.)

NOTES:
1. In criterion 1, the phrase “one or more blood cultures” means that at least one bottle from a blood draw is reported by the laboratory as having grown organisms (i.e., is a positive blood culture).
2. In criterion 1, the term “recognized pathogen” does not include organisms considered common commensals (see criteria 2 and 3 for a list of common commensals). A few of the recognized pathogens are S. aureus, Enterococcus spp., E. coli, Pseudomonas spp., Klebsiella spp., Candida spp., etc.
3. In criteria 2 and 3, the phrase “two or more blood cultures drawn on separate occasions” means 1) that blood from at least two blood draws were collected within two days of each other (e.g., blood draws on Monday and Tuesday or Monday and Wednesday would be acceptable for blood cultures drawn on separate occasions, but blood draws on Monday and Thursday would be too far apart in time to meet this criterion), and 2) that at least one bottle from each blood draw is reported by the laboratory as having grown the same common commensal (i.e., is a positive blood culture). (See Note 4 for determining sameness of organisms.)
   a. For example, an adult patient has blood drawn at 8 a.m. and again at 8:15 a.m. of the same day. Blood from each blood draw is inoculated into two bottles and incubated (four bottles total). If one bottle from each blood draw set is positive for coagulase-negative staphylococci, this part of the criterion is met.
   b. For example, a neonate has blood drawn for culture on Tuesday and again on Saturday and both grow the same common commensal. Because the time between these blood cultures exceeds the two-day period for blood draws stipulated in criteria 2 and 3, this part of the criteria is not met.
   c. A blood culture may consist of a single bottle for a pediatric blood draw due to volume constraints. Therefore, to meet this part of the criterion, each bottle from two or more draws would have to be culture-positive for the same commensal.
4. If the common commensal is identified to the species level from one culture, and a companion culture is identified with only a descriptive name (e.g., to the genus level), then it is assumed that the organisms are the same. The organism identified to the species level should be reported as the infecting pathogen along with its antibiogram if available (see Table 1 below).
Table 1. Examples of how to report speciated and unspeciated common commensals

<table>
<thead>
<tr>
<th>Culture Report</th>
<th>Companion Culture Report</th>
<th>Report as…</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>S. epidermidis</em></td>
<td>Coagulase-negative staphylococci</td>
<td><em>S. epidermidis</em></td>
</tr>
<tr>
<td><em>Bacillus</em> spp. (not <em>anthracis</em>)</td>
<td><em>B. cereus</em></td>
<td><em>B. cereus</em></td>
</tr>
<tr>
<td><em>S. salivarius</em></td>
<td>Strep viridans</td>
<td><em>S. salivarius</em></td>
</tr>
</tbody>
</table>

5. Only genus and species identification should be utilized to determine the sameness of organisms. No additional comparative methods should be used (e.g., morphology or antibiograms) because laboratory testing capabilities and protocols may vary between facilities. This will reduce reporting variability, solely due to laboratory practice, between facilities reporting LCBIs meeting criterion 2. Report the organism to the genus/species level only once, and if antibiogram data are available, report the results from the most resistant panel.

6. LCBI criteria 1 and 2 may be used for patients of any age, including patients ≤ 1 year of age.

7. Specimen Collection Considerations:
   Ideally, blood specimens for culture should be obtained from two to four blood draws from separate venipuncture sites (e.g., right and left antecubital veins), not through a vascular catheter. These blood draws should be performed simultaneously or over a short period of time (i.e., within a few hours). If your facility does not currently obtain specimens using this technique, you must still report BSIs using the criteria and notes above, but you should work with appropriate personnel to facilitate better specimen collection practices for blood cultures.

REPORTING INSTRUCTIONS:
- Report organisms cultured from blood as BSI – LCBI when no other site of infection is evident.
- When there is a positive blood culture and clinical signs or symptoms of localized infection at a vascular access site, but no other infection can be found, the infection is considered a primary BSI.
- Purulent phlebitis confirmed with a positive semiquantitative culture of a catheter tip, but with either negative or no blood culture is considered a CVS-VASC, not a BSI nor an SST-SKIN or ST infection.
- Occasionally a patient with both peripheral and central IV lines develops a primary bloodstream infection (LCBI) that can clearly be attributed to the peripheral line (e.g., pus at the insertion site and matching pathogen from pus and blood). In this situation, enter “Central Line = No” in the NHSN application. You should, however, include the patient’s central line days in the summary denominator count.

June, 2011 4-5
**Numerator Data:** The *Primary Bloodstream Infection (BSI)* form (CDC 57.108) is used to collect and report each CLABSI that is identified during the month selected for surveillance. The *Instructions for Completion of Primary Bloodstream Infection Form* (Tables of Instructions, Tables 2 and 2a.) contains brief instructions for collection and entry of each data element on the form. The Primary BSI form includes patient demographic information and whether a central line was present, and, if so, the type of central line the patient had as appropriate to the location; these data will be used to calculate line-specific infection rates. Additional data include the specific criteria met for identifying the primary BSI, whether the patient died, the organisms isolated from blood cultures, and the organisms’ antimicrobial susceptibilities.

**Denominator Data:** Device days and patient days are used for denominators (see Chapter 16, Key Terms). Device-day denominator data that are collected differ according to the location of the patients being monitored; however, they should be collected at the same time each day. When denominator data are available from electronic databases, these sources may be used as long as the counts are not substantially different (+/- 5%) from manually-collected counts.

For ICUs and locations other than specialty care areas (SCAs) and NICUs, the number of patients with one or more central lines of any type is collected daily, at the same time each day, during the month and recorded on the *Denominators for Intensive Care Unit (ICU)/Other Locations (Not NICU or Specialty Care Area (SCA))* (CDC 57.118). Only the totals for the month are entered into NHSN. When denominator data are available from electronic sources (e.g., central line days from electronic charting), these sources may be used as long as the counts are not substantially different (+/- 5%) from manually-collected counts.

For specialty care areas, the number of patients with one or more central lines is dichotomized into those with permanent central lines and those with temporary central lines on the *Denominators for Specialty Care Area* (CDC 57.117) form. Each is collected daily, at the same time each day. Only the total for the month are entered into NHSN. This distinction in lines is made because permanent lines are commonly used in patients frequenting these areas and may have lower rates of associated infection than central lines inserted for temporary use. If a patient has both a temporary and a permanent central line, count the day only as a temporary line day. The *Instructions for Completion of Denominators for Intensive Care Unit (ICU)/Other Locations Form* (Tables of Instructions, Table 6) and *Instructions for Completion of Denominators for Specialty Care Areas (SCA)* Form (Tables of Instructions, Table 7) contain brief instructions for collection and entry of each data element on the forms.

In NICUs, again because of differing infection risks, the number of patients with central lines and those with umbilical catheters is collected daily, at the same time each day, during the month. If a patient has both an umbilical catheter and a central line, count the day only as an umbilical catheter day. On the *Denominators for Neonatal Intensive Care Form*...
Unit (NICU) (CDC 57.116) form, patients are further stratified by birthweight in five categories since risk of BSI also varies by birthweight.

NOTE: The weight of the infant at the time of BSI is not used and should not be reported. For example, if a neonate weighs 1006 grams at birth but remains in the NICU for two months and has a body weight of 1650 grams when it develops a CLABSI, record the birthweight of 1006 grams on the BSI form. The Instructions for Completion of Denominators for Neonatal Intensive Care Unit (NICU) form (Tables of Instructions, Table 8) contains brief instructions for collection and entry of each data element on the forms.

Data Analyses: The SIR is calculated by dividing the number of observed infections by the number of expected infections. The number of expected infections, in the context of statistical prediction, is calculated using CLABSI rates from a standard population during a baseline time period as reported in the NHSN Report.

NOTE: The SIR will be calculated only if the number of expected HAIs (numExp) is ≥ 1.

\[
\text{SIR} = \frac{\text{Observed (O) HAIs}}{\text{Expected (E) HAIs}}
\]

While the CLABSI SIR can be calculated for single locations, the measure also allows you to summarize your data across multiple locations, adjusting for differences in the incidence of infection among the location types. For example, you will be able to obtain one CLABSI SIR adjusting for all locations reported. Similarly, you can obtain one CLABSI SIR for all specialty care areas in your facility.

The CLABSI rate per 1000 central line days is calculated by dividing the number of CLABSI by the number of central line days and multiplying the result by 1000. The Central Line Utilization Ratio is calculated by dividing the number of central line days by the number of patient days. These calculations will be performed separately for different types of ICUs, specialty care areas, and other locations in the institution. Separate rates and ratios will also be calculated for different types of catheters in specialty care areas and NICUs, and for birthweight categories in NICUs.


CLABS Criteria 1 & 2 – Used for Patients of any Age

Blood Cultures (list here)
- Date: _____ Organism: ______ #_______
- Date: _____ Organism: ______ #_______
- Date: _____ Organism: ______ #_______
- Date: _____ Organism: ______ #_______
- Date: _____ Organism: ______ #_______
- Date: _____ Organism: ______ #_______
- Date: _____ Organism: ______ #_______
- Date: _____ Organism: ______ #_______

Non-blood cultures (list here)
- Date: _____ Source: ___ Organism: ______
- Date: _____ Source: ___ Organism: ______
- Date: _____ Source: ___ Organism: ______
- Date: _____ Source: ___ Organism: ______
- Date: _____ Source: ___ Organism: ______
- Date: _____ Source: ___ Organism: ______
- Date: _____ Source: ___ Organism: ______
- Date: _____ Source: ___ Organism: ______

Patient has a recognized pathogen cultured from one or more blood cultures (circle pathogen in blood culture section & check yes)

Organism cultured from blood is not related to an infection at another site (check if not related)

Yes (not related) – Meets Criteria

CIRCLE CONCLUSION
YES
NO
INDETERMINATE

Common commensal cultured from two or more blood cultures drawn on separate occasions (circle commensals in blood culture section & check yes)

Organism cultured from blood is not related to an infection at another site (check if not related)

No – Does not meet NHSN

Patient has at least one of the following signs or symptoms (circle all that apply): (check if yes)
- Fever (>38 C or 100.4 F)
- Chills
- Hypotension

Admit Temp: _________
Admit Blood Pressure: ______

Date: _____ Symptom: _________
Date: _____ Symptom: _________
Date: _____ Symptom: _________

Signs or symptoms not related to an infection at another site (check if not related)

Yes (not related) – Meets Criteria 2

No – Does not meet NHSN

Notes: Dates & site notes if needed: ____________________________________________________________________
Central Line- Associated Bloodstream Infection (CLABSI) Event Check List Criteria

Instructions for use

Use this check list to determine if an infection meets the NHSN CLABSI criteria.

Blood cultures: Fill in relevant blood cultures in blood culture section, if you need to ensure there are 2 different blood cultures then include the unique identifying number for the blood culture next to #______________.

Non-blood cultures: Fill in the relevant cultures to help you determine if a secondary infection exists.

Then go down the flow sheet using Criteria 1 if you find a recognized pathogen and Criteria 2 if you find common commensals cultured from 2 or more blood cultures. As instructed in the flow sheet, circle the relevant cultures used to assist in make the CLABSI determination, for example, you would circle the recognized pathogen in the blood cultures box that was used to come up with a determination of a CLABSI. If none, do not circle.

You must understand the terms defined in bold below to complete the criteria checklist. You will find the terms in bold letters in the checklist.

Central line defined: An intravascular catheter (used for infusion, withdrawal of blood, or hemodynamic monitoring) that terminates at or close to the heart or in one of the great vessels which includes the: aorta, pulmonary artery, superior vena cava, inferior vena cava, brachiocephalic veins, internal jugular veins, subclavian veins, external iliac veins, common iliac veins, femoral veins, and in neonates, the umbilical artery/vein.

The Central line (CL) must be in place or within 48 hours of CL discontinuation when blood cultures drawn are drawn.

Common Commensals include but are not limited to (for a more extensive list, please refer to common commensal list):

- Diphtheroids (Corynebacterium spp. not C. diphtheria
- Bacillus spp. not B. anthracis, Propionibacterium spp.
- Coagulase-negative staphylococci including S. epidermidis
- Viridans group streptococci, Aerococcus spp., Micococcus spp., S. salivarius

Recognized pathogens (Do Not include common commensals) include but are not limited to:

- S. aureus
- Enterococcus spp.
- E. coli
- Pseudomonas spp.
- Klebsiella spp.
- Candida spp.

One or more blood cultures means at least one bottle from each blood draw (each draw/culture requires 2 bottles) is reported by the laboratory to have grown organisms. It is a positive blood culture. For example, two blood cultures would require 4 bottles and for both blood cultures to be positive, one bottle from each set would have to grown organisms.

Two or more blood cultures drawn on separate occasions means:

1. Blood from at least 2 blood draws were collected within two days of each other
2. At least one bottle from each blood draw is reported by the laboratory as having grown the same common commensal. It is considered a positive blood culture.

Pediatric blood draw consideration

Blood culture may consist of a single bottle for a pediatric blood draw. Therefore to meet criteria, each bottle from two or more draws would have to be culture-positive for the same commensal.
Infection at another site:

In Criteria’s 1 or 2 in order to make the determination of a CLABS you must ensure that the organism cultured from the blood is not related to an infection at another site. Look at non-blood culture results to see if the organism cultured in the blood is the same as an organism cultured from a different source. In addition, refer to NHSN manual, Chapter 17 to see if the signs or symptoms a patient is having meet the NHSN criteria for an infection. If so, the organism cultured from the blood would be related to an infection at another site and would not meet NHSN criteria for a CLABS.

Other Considerations:

Patient has a peripheral IV and central line (CL) in place at the same time:

Primary BSI attributed to peripheral line and not the central line if pus at the peripheral line insertion site matches the blood pathogen.

Catheter tip cultures

Purulent phlebitis confirmed with a positive semiquantitative culture of a catheter tip, but with either negative or no blood culture is considered a cardiovascular system-venous arterial system related infection, not a BSI or CLBSI.

Localized infection at Central Line site:

A positive blood culture and localized infection at the central line site and no other infection would be considered a primary BSI.

For further details refer to the Device-associated CLBSI module
### VAP

<table>
<thead>
<tr>
<th>event category</th>
<th>full term</th>
<th>short term</th>
<th>specifications</th>
<th>standard</th>
</tr>
</thead>
<tbody>
<tr>
<td>healthcare-associated infection</td>
<td>pneumonia that is ventilator-associated</td>
<td>VAP</td>
<td></td>
<td>AHRQ, NHSN</td>
</tr>
</tbody>
</table>

**implementation guidance**

NHSN guidance included on the next few pages.

**examples**
Ventilator-Associated Pneumonia (VAP) Event

**Introduction:** In 2002, an estimated 250,000 healthcare-associated pneumonias developed in U.S. hospitals and 36,000 of these were associated with deaths. Patients with mechanically-assisted ventilation have a high risk of developing healthcare-associated pneumonia. From 2006-2007, within NHSN facilities almost 5,400 VAPs were reported and incidence for various types of hospital units ranged from 2.1-11.0 per 1,000 ventilator days.


**Settings:** Surveillance will occur in any inpatient location where denominator data can be collected, which may include critical/intensive care units (ICU), specialty care areas (SCA), neonatal units, including neonatal intensive care units (NICUs), stepdown units, wards, and long term care units. A complete listing of inpatient locations can be found in Chapter 15.

NOTE: It is not required to monitor for VAPs after the patient is discharged from the facility, however, if discovered, a VAP should be reported to NHSN. No additional ventilator days are reported.

**Requirements:** Surveillance for VAP in at least one inpatient location in the healthcare institution for at least one calendar month as indicated in the *Patient Safety Monthly Reporting Plan* (CDC 57.106).

**Definitions:** As for all infections reported to NHSN, infections associated with complications or extensions of infections already present on admission, unless a change in pathogen or symptoms strongly suggests the acquisition of a new infection area not considered healthcare associated. Therefore, infections that become apparent within the first few days of admission must be carefully reviewed to determine whether they should be considered healthcare associated.

Pneumonia (PNEU) is identified by using a combination of radiologic, clinical and laboratory criteria. The following pages outline the various assessment criteria that may be used for meeting the surveillance definition of healthcare-associated pneumonia (Tables 2-5 and Figures 1 and 2). Report PNEUs that are ventilator-associated (i.e., patient was intubated and ventilated at the time of, or within 48 hours before, the onset of the event).

NOTE: There is no minimum period of time that the ventilator must be in place in order for the PNEU to be considered ventilator associated.

**Location of attribution:** The inpatient location where the patient was assigned on the date of the PNEU event, which is further defined as the date when the first clinical evidence appeared or the date the specimen used to meet the PNEU criterion was collected, whichever came first.

EXAMPLE: Patient is intubated and ventilated in the Operating Room and then is admitted to the MICU. Within 24 hours of admission to the MICU, patient meets criteria for PNEU. This is
reported to NHSN as a VAP for the MICU, because the Operating Room is not an inpatient location and no denominator data are collected there.

TRANSFER RULE EXCEPTION: If a VAP develops within 48 hours of transfer from one inpatient location to another in the same facility or a new facility, the infection is attributed to the transferring location. This is called the Transfer Rule and examples are shown below:

- Patient on a ventilator in the SICU is transferred to the surgical ward. Thirty six (36) hours later, the patient meets the criteria for PNEU. This is reported to NHSN as a VAP for the SICU.
- Patient is transferred to the medical ward from the MSICU after having ventilator removed. Within 24 hours, the patient meets criteria for a PNEU. This is reported to NHSN as a VAP for the MSICU.
- Patient on a ventilator is transferred from the medical ward to the coronary care ICU (CCU). After 4 days in the CCU, the patient meets the criteria for a PNEU. This is reported to NHSN as a VAP for the CCU.
- Patient on the Respiratory ICU (RICU) of Hospital A had the endotracheal tube and ventilator removed and is discharged home a few hours later. The ICP from Hospital B calls the next day to report that this patient has been admitted to Hospital B with a PNEU. This VAP should be reported to NHSN for, and by, Hospital A and attributed to the RICU. No additional ventilator days are reported.

Ventilator: A device to assist or control respiration continuously, inclusive of the weaning period, through a tracheostomy or by endotracheal intubation.

NOTE: Lung expansion devices such as intermittent positive-pressure breathing (IPPB); nasal positive end-expiratory pressure (PEEP); and continuous nasal positive airway pressure (CPAP, hypoCPAP) are not considered ventilators unless delivered via tracheostomy or endotracheal intubation (e.g., ET-CPAP).

General Comments Applicable to All Pneumonia Specific Site Criteria:

1. Physician’s diagnosis of pneumonia alone is not an acceptable criterion for healthcare-associated pneumonia.
2. Although specific criteria are included for infants and children, pediatric patients may meet any of the other pneumonia specific site criteria.
3. Ventilator-associated pneumonia (i.e., pneumonia in persons who had a device to assist or control respiration continuously through a tracheostomy or by endotracheal intubation within the 48-hour period before the onset of infection, inclusive of the weaning period) should be so designated when reporting data.
4. When assessing a patient for presence of pneumonia, it is important to distinguish between changes in clinical status due to other conditions such as myocardial infarction, pulmonary embolism, respiratory distress syndrome, atelectasis, malignancy, chronic obstructive pulmonary disease, hyaline membrane disease, bronchopulmonary dysplasia, etc. Also, care must be taken when assessing intubated patients to distinguish between tracheal colonization, upper respiratory tract infections (e.g., tracheobronchitis), and early onset pneumonia. Finally, it should be recognized that it may be difficult to determine healthcare-associated pneumonia in the elderly, infants, and immunocompromised patients since such conditions
may mask typical signs or symptoms associated with pneumonia. Alternate specific criteria for the elderly, infants and immunocompromised patients have been included in this definition of healthcare-associated pneumonia.

5. Healthcare-associated pneumonia can be characterized by its onset: early or late. Early onset pneumonia occurs during the first four days of hospitalization and is often caused by *Moraxella catarrhalis, H. influenzae,* and *S. pneumoniae.* Causative agents of late onset pneumonia are frequently gram negative bacilli or *S. aureus,* including methicillin-resistant *S. aureus.* Viruses (e.g., Influenza A and B or Respiratory Syncytial Virus) can cause early and late onset healthcare-associated pneumonia, whereas yeasts, fungi, legionellae, and *Pneumocystis carinii* are usually pathogens of late onset pneumonia.

6. Pneumonia due to gross aspiration (for example, in the setting of intubation in the emergency room or operating room) is considered healthcare-associated if it meets any specific criteria and was not clearly present or incubating at the time of admission to the hospital.

7. Multiple episodes of healthcare-associated pneumonia may occur in critically ill patients with lengthy hospital stays. When determining whether to report multiple episodes of healthcare-associated pneumonia in a single patient, look for evidence of resolution of the initial infection. The addition of or change in pathogen alone is not indicative of a new episode of pneumonia. The combination of new signs and symptoms and radiographic evidence or other diagnostic testing is required.

8. Positive Gram stain for bacteria and positive KOH (potassium hydroxide) mount for elastin fibers and/or fungal hyphae from appropriately collected sputum specimens are important clues that point toward the etiology of the infection. However, sputum samples are frequently contaminated with airway colonizers and therefore must be interpreted cautiously. In particular, *Candida* is commonly seen on stain, but infrequently causes healthcare-associated pneumonia.

**Table 1: Abbreviations used in PNEU laboratory criteria**

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>BAL</td>
<td>bronchoalveolar lavage</td>
</tr>
<tr>
<td>EIA</td>
<td>enzyme immunoassay</td>
</tr>
<tr>
<td>FAMA</td>
<td>fluorescent-antibody staining of membrane antigen</td>
</tr>
<tr>
<td>IFA</td>
<td>immunofluorescent antibody</td>
</tr>
<tr>
<td>LRT</td>
<td>lower respiratory tract</td>
</tr>
<tr>
<td>PCR</td>
<td>polymerase chain reaction</td>
</tr>
<tr>
<td>PMN</td>
<td>polymorphonuclear leukocyte</td>
</tr>
<tr>
<td>RIA</td>
<td>radioimmunoassay</td>
</tr>
</tbody>
</table>

**REPORTING INSTRUCTIONS:**

- There is a hierarchy of specific categories within the major site pneumonia. Even if a patient meets criteria for more than one specific site, report only one:
  - If a patient meets criteria for both PNU1 and PNU2, report PNU2
  - If a patient meets criteria for both PNU2 and PNU3, report PNU3
  - If a patient meets criteria for both PNU1 and PNU3, report PNU3
- Report concurrent lower respiratory tract infection (e.g., abscess or empyema) and pneumonia with the same organism(s) as pneumonia
- Lung abscess or empyema without pneumonia are classified as LUNG
- Bronchitis, tracheitis, tracheobronchitis, or bronchiolitis without pneumonia are classified as BRON.

**Table 2: Specific Site Algorithms for Clinically Defined Pneumonia (PNU1)**
### Radiology

- Two or more serial chest radiographs with at least one of the following:
  - New or progressive and persistent infiltrate
  - Consolidation
  - Cavitation
  - Pneumatoceles, in infants ≤ 1 year old

### Signs/Symptoms/Laboratory

FOR ANY PATIENT, at least one of the following:

- Fever (>38°C or >100.4°F) with no other recognized cause
- Leukopenia (<4000 WBC/mm³) or leukocytosis (≥12,000 WBC/mm³)
- For adults ≥ 70 years old, altered mental status with no other recognized cause

and at least two of the following:

- New onset of purulent sputum³, or change in character of sputum⁴, or increased respiratory secretions, or increased suctioning requirements
- New onset or worsening cough, or dyspnea, or tachypnea⁵
- Rales⁶ or bronchial breath sounds
- Worsening gas exchange (e.g. O₂ desaturations [e.g. pulse oximetry < 94%], increased oxygen requirements, or increased ventilator demand)

ALTERNATE CRITERIA, for infants ≤ 1 year old:

Worsening gas exchange (e.g. O₂ desaturations [e.g. pulse oximetry < 94%], increased oxygen requirements, or increased ventilator demand)

and at least three of the following:

- Temperature instability with no other recognized cause
- Leukopenia (<4000 WBC/mm³) or leukocytosis (≥15,000 WBC/mm³) and left shift (>10% band forms)
- New onset of purulent sputum³ or change in character of sputum⁴, or increased respiratory secretions or increased suctioning requirements
- Apnea, tachypnea⁵, nasal flaring with retraction of chest wall or grunting
- Wheezing, rales⁶, or rhonchi
- Cough
- Bradycardia (<100 beats/min) or tachycardia (>170 beats/min)

ALTERNATE CRITERIA, for child > 1 year old or ≤ 12 years old, at least three of the following:

- Fever (>38.4°C or >101.1°F) or hypothermia (<36.5°C or <97.7°F) with no other recognized cause
- Leukopenia (<4000 WBC/mm³) or leukocytosis (≥15,000 WBC/mm³)
- New onset of purulent sputum³, or change in character of sputum⁴, or increased respiratory secretions, or increased suctioning requirements
- New onset or worsening cough, or dyspnea, apnea, or tachypnea⁵.
- Rales⁶ or bronchial breath sounds.
- Worsening gas exchange (e.g. O₂ desaturations [e.g. pulse oximetry < 94%], increased oxygen requirements, or increased ventilator demand)
### Table 3: Specific Site Algorithms for Pneumonia with Common Bacterial or Filamentous Fungal Pathogens and Specific Laboratory Findings (PNU2)

<table>
<thead>
<tr>
<th>Radiology</th>
<th>Signs/Symptoms</th>
<th>Laboratory</th>
</tr>
</thead>
<tbody>
<tr>
<td>Two or more serial chest radiographs with at least one of the following(^{1,2}):</td>
<td>At least one of the following:</td>
<td>At least one of the following:</td>
</tr>
<tr>
<td>New or progressive and persistent infiltrate</td>
<td>Fever ((&gt;38^\circ)C or (&gt;100.4^\circ)F) with no other recognized cause</td>
<td>Positive growth in blood culture(^8) not related to another source of infection</td>
</tr>
<tr>
<td>Consolidation</td>
<td>Leukopenia ((&lt;4000) WBC/mm(^3)) or leukocytosis ((\geq 12,000) WBC/mm(^3))</td>
<td>Positive growth in culture of pleural fluid</td>
</tr>
<tr>
<td>Cavitation</td>
<td>For adults (\geq 70) years old, altered mental status with no other recognized cause</td>
<td>Positive quantitative culture(^9) from minimally contaminated LRT specimen (e.g., BAL or protected specimen brushing)</td>
</tr>
<tr>
<td>Pneumatoceles, in infants (\leq 1) year old and at least one of the following:</td>
<td></td>
<td>(\geq 5%) BAL-obtained cells contain intracellular bacteria on direct microscopic exam (e.g., Gram stain)</td>
</tr>
<tr>
<td>NOTE: In patients without underlying pulmonary or cardiac disease (e.g., respiratory distress syndrome, bronchopulmonary dysplasia, pulmonary edema, or chronic obstructive pulmonary disease), one definitive chest radiograph is acceptable.(^1)</td>
<td>New onset of purulent sputum(^3), or change in character of sputum(^5), or increased respiratory secretions, or increased suctioning requirements</td>
<td>Histopathologic exam shows at least one of the following evidences of pneumonia:</td>
</tr>
<tr>
<td></td>
<td>New onset or worsening cough, or dyspnea or tachypnea(^5)</td>
<td>Abscess formation or foci of consolidation with intense PMN accumulation in bronchioles and alveoli</td>
</tr>
<tr>
<td></td>
<td>Rales(^6) or bronchial breath sounds</td>
<td>Positive quantitative culture(^9) of lung parenchyma</td>
</tr>
<tr>
<td></td>
<td>Worsening gas exchange (e.g., (O_2) desaturations [e.g., (PaO_2/FiO_2 \leq 240)(^1), increased oxygen requirements, or increased ventilator demand)</td>
<td>Evidence of lung parenchyma invasion by fungal hyphae or pseudohyphae</td>
</tr>
</tbody>
</table>
Table 4: Specific Site Algorithms for Viral, Legionella, and other Bacterial Pneumonias with Definitive Laboratory Findings (PNU2)

<table>
<thead>
<tr>
<th>Radiology</th>
<th>Signs/Symptoms</th>
<th>Laboratory</th>
</tr>
</thead>
<tbody>
<tr>
<td>Two or more serial chest radiographs with at least one of the following:\nNew or progressive and persistent infiltrate\nConsolidation\nCavitation\nPneumatoceles, in infants ≤ 1 year old</td>
<td>At least one of the following:\nFever (&gt;38°C or &gt;100.4°F) with no other recognized cause\nLeukopenia (&lt;4000 WBC/mm³) or leukocytosis (≥12,000 WBC/mm³)\nFor adults ≥70 years old, altered mental status with no other recognized cause and at least one of the following:\nNew onset of purulent sputum³, or change in character of sputum⁴, or increased respiratory secretions, or increased suctioning requirements\nNew onset or worsening cough or dyspnea, or tachypnea⁵\nRales⁶ or bronchial breath sounds\nWorsening gas exchange (e.g. O₂ desaturations [e.g., PaO₂/FIO₂ &lt; 240], increased oxygen requirements, or increased ventilator demand)</td>
<td>At least one of the following\nPositive culture of virus or Chlamydia from respiratory secretions\nPositive detection of viral antigen or antibody from respiratory secretions (e.g., EIA, FAMA, shell vial assay, PCR)\nFourfold rise in paired sera (IgG) for pathogen (e.g., influenza viruses, Chlamydia)\nPositive PCR for Chlamydia or Mycoplasma\nPositive micro-IF test for Chlamydia\nPositive culture or visualization by micro-IF of Legionella spp, from respiratory secretions or tissue.\nDetection of Legionella pneumophila serogroup 1 antigens in urine by RIA or EIA\nFourfold rise in L. pneumophila serogroup 1 antibody titer to ≥1:128 in paired acute and convalescent sera by indirect IFA.</td>
</tr>
</tbody>
</table>
Table 5: Specific Site Algorithm for Pneumonia in Immunocompromised Patients (PNU3)

<table>
<thead>
<tr>
<th>Radiology</th>
<th>Signs/Symptoms</th>
<th>Laboratory</th>
</tr>
</thead>
<tbody>
<tr>
<td>Two or more serial chest radiographs with at least one of the following:&lt;br&gt; New or progressive and persistent infiltrate&lt;br&gt; Consolidation&lt;br&gt; Cavitation&lt;br&gt; Pneumatoceles, in infants ≤ 1 year old</td>
<td>Patient who is immunocompromised (^{13}) has at least one of the following:&lt;br&gt; Fever (&gt;38°C or &gt;100.4°F) with no other recognized cause&lt;br&gt; For adults ≥70 years old, altered mental status with no other recognized cause&lt;br&gt; New onset of purulent sputum (^{2}), or change in character of sputum (^{4}), or increased respiratory secretions, or increased suctioning requirements&lt;br&gt; New onset or worsening cough, or dyspnea, or tachypnea (^{3})&lt;br&gt; Rales (^{6}) or bronchial breath sounds&lt;br&gt; Worsening gas exchange (e.g. (\text{O}_2) desaturations [e.g., (\text{PaO}_2/\text{FiO}_2 \leq 240)](^{7}), increased oxygen requirements, or increased ventilator demand)&lt;br&gt; Hemoptysis&lt;br&gt; Pleuritic chest pain</td>
<td>At least one of the following:&lt;br&gt; Matching positive blood and sputum cultures with (Candida) spp. (^{14,15})&lt;br&gt; Evidence of fungi or (Pneumocystis carinii) from minimally contaminated LRT specimen (e.g., BAL or protected specimen brushing) from one of the following:&lt;br&gt; - Direct microscopic exam&lt;br&gt; - Positive culture of fungi&lt;br&gt; Any of the following from LABORATORY CRITERIA DEFINED UNDER PNU2</td>
</tr>
</tbody>
</table>

NOTE: In patients without underlying pulmonary or cardiac disease (e.g. respiratory distress syndrome, bronchopulmonary dysplasia, pulmonary edema, or chronic obstructive pulmonary disease), one definitive chest radiograph is acceptable.\(^{3}\)

Footnotes to Algorithms:
1. Occasionally, in nonventilated patients, the diagnosis of healthcare-associated pneumonia may be quite clear on the basis of symptoms, signs, and a single definitive chest radiograph. However, in patients with pulmonary or cardiac disease (for example, interstitial lung disease or congestive heart failure), the diagnosis of pneumonia may be particularly difficult. Other non-infectious conditions (for example, pulmonary edema from decompensated congestive heart failure) may simulate the presentation of pneumonia. In these more difficult cases, serial chest radiographs must be examined to help separate infectious from non-infectious pulmonary processes. To help confirm difficult cases, it may be useful to review radiographs on the day of diagnosis, 3 days prior to the diagnosis and on days 2 and 7 after the diagnosis. Pneumonia may have rapid onset and progression, but does not resolve quickly. Radiographic changes of pneumonia persist for several weeks. As a result, rapid radiographic resolution suggests that the patient does not have pneumonia, but rather a non-infectious process such as atelectasis or congestive heart failure.
2. Note that there are many ways of describing the radiographic appearance of pneumonia. Examples include, but are not limited to, “air-space disease”, “focal opacification”, “patchy areas of increased density”. Although perhaps not specifically delineated as pneumonia by the radiologist, in the appropriate clinical setting these alternative descriptive wordings should be seriously considered as potentially positive findings.
3. Purulent sputum is defined as secretions from the lungs, bronchi, or trachea that contain ≥25 neutrophils and ≤10 squamous epithelial cells per low power field (x100). If your laboratory reports these data qualitatively (e.g., “many WBCs” or “few squames”), be sure their descriptors match this definition of purulent sputum. This laboratory confirmation is required since written clinical descriptions of purulence are highly variable.

4. A single notation of either purulent sputum or change in character of the sputum, is not meaningful; repeated notations over a 24 hour period would be more indicative of the onset of an infectious process. Change in character of sputum refers to the color, consistency, odor and quantity.

5. In adults, tachypnea is defined as respiration rate >25 breaths per minute. Tachypnea is defined as >75 breaths per minute in premature infants born at <37 weeks gestation and until the 40th week; >60 breaths per minute in patients <2 months old; >50 breaths per minute in patients 2-12 months old; and >30 breaths per minute in children >1 year old.

6. Rales may be described as “crackles”.

7. This measure of arterial oxygenation is defined as the ratio of the arterial tension (PaO₂) to the inspiratory fraction of oxygen (FiO₂).

8. Care must be taken to determine the etiology of pneumonia in a patient with positive blood cultures and radiographic evidence of pneumonia, especially if the patient has invasive devices in place such as intravascular lines or an indwelling urinary catheter. In general, in an immunocompetent patient, blood cultures positive for coagulase negative staphylococci, common skin contaminants, and yeasts will not be the etiologic agent of the pneumonia.

9. Refer to Threshold values for cultured specimens (Table 6). An endotracheal aspirate is not a minimally contaminated specimen. Therefore, an endotracheal aspirate does not meet the laboratory criteria.

10. Once laboratory-confirmed cases of pneumonia due to respiratory syncytial virus (RSV), adenovirus, or influenza virus have been identified in a hospital, clinician’s presumptive diagnosis of these pathogens in subsequent cases with similar clinical signs and symptoms is an acceptable criterion for presence of healthcare-associated infection.

11. Scant or watery sputum is commonly seen in adults with pneumonia due to viruses and Mycoplasma although sometimes the sputum may be mucopurulent. In infants, pneumonia due to RSV or influenza yields copious sputum. Patients, except premature infants, with viral or mycoplasmal pneumonia may exhibit few signs or symptoms, even when significant infiltrates are present on radiographic exam.

12. Few bacteria may be seen on stains of respiratory secretions from patients with pneumonia due to Legionella spp, mycoplasma, or viruses.

13. Immunocompromised patients include those with neutropenia (absolute neutrophil count <500/mm³), leukemia, lymphoma, HIV with CD4 count <200, or splenectomy; those who are early post-transplant, are on cytotoxic chemotherapy, or are on high dose steroids (e.g., >40mg of prednisone or its equivalent (>160mg hydrocortisone, >32mg methylprednisolone, >6mg dexamethasone, >200mg cortisone) daily for >2weeks).

14. Blood and sputum specimens must be collected within 48 hours of each other.

15. Semiquantitative or nonquantitative cultures of sputum obtained by deep cough, induction, aspiration, or lavage are acceptable. If quantitative culture results are available, refer to algorithms that include such specific laboratory findings.
Figure 2: Pneumonia Flow Diagram, Alternative Criteria for Infants and Children

**PNEUMONIA FLOW DIAGRAM**
**ALTERNATE CRITERIA FOR INFANTS AND CHILDREN**

**Facility ID # _____________**
**Event # _____________**
**Event Date __/__/_________**

**Instructions: Complete form only if x-ray criteria are met**

- **Infants < 1 y.o.**
  - Worsening gas exchange (e.g., O₂ desats [e.g., pulse oximetry <94%], ↑ O₂ req, or ↑ ventilation demand)
  - Temperature instability with no other recognized cause
  - Leukopenia (< 4,000 WBC/mm³) or leukocytosis (> 15,000 WBC/mm³) and left shift (> 10% band forms)
  - New onset of purulent sputum, or change in character of sputum, or ↑ respiratory secretions, or ↑ suctioning requirements
  - Apnea, tachypnea, nasal flaring with retraction of chest wall or grunting
  - Wheezing, rales, or rhonchi
  - Cough
  - Bradycardia (<100 beats/min.) or tachycardia (> 170 beats/min.)

- **Patient with underlying diseases** has 2 or more serial X-rays with one of the following:
  - New or progressive and persistent infiltrate
  - Consolidation
  - Cavitation
  - Pneumatoceles, in ≤ 1 y.o.

- **Children >1 or < 12 y.o.**
  - Fever (>38.4°C/101.1°F) or hypothermia (< 36.5°C/97.7°F) with no other recognized cause
  - Leukopenia (< 4,000 WBC/mm³) or leukocytosis (> 15,000 WBC/mm³)
  - New onset of purulent sputum, or change in character of sputum, or ↑ respiratory secretions, or ↑ suctioning requirements
  - Apnea, tachypnea, nasal flaring with retraction of chest wall or grunting
  - Wheezing, rales, or rhonchi
  - Cough
  - Bradycardia (<100 beats/min.) or tachycardia (> 170 beats/min.)

- **Patient without underlying diseases** has 1 or more serial X-rays with one of the following:
  - New or progressive and persistent infiltrate
  - Consolidation
  - Cavitation
  - Pneumatoceles, in ≤ 1 y.o.

- **Children >1 or < 12 y.o.**
  - At least three of the following:
    - Fever (>38.4°C/101.1°F) or hypothermia (< 36.5°C/97.7°F) with no other recognized cause
    - Leukopenia (< 4,000 WBC/mm³) or leukocytosis (> 15,000 WBC/mm³)
    - New onset of purulent sputum, or change in character of sputum, or ↑ respiratory secretions, or ↑ suctioning requirements
    - New onset or worsening cough, or dyspnea, apnea, or tachypnea
    - Rales or bronchial breath sounds
    - Worsening gas exchange (e.g., O₂ desats [e.g., pulse oximetry < 94%], ↑ O₂ req, or ↑ ventilation demand)

- **Patient with underlying diseases** has 2 or more serial X-rays with one of the following:
  - New or progressive and persistent infiltrate
  - Consolidation
  - Cavitation
  - Pneumatoceles, in ≤ 1 y.o.

**PNU1:**
**Clinically defined pneumonia**
**Table 6: Threshold values for cultured specimens used in the diagnosis of pneumonia**

<table>
<thead>
<tr>
<th>Specimen collection/technique</th>
<th>Values</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lung parenchyma*</td>
<td>≥10⁴ cfu/g tissue</td>
</tr>
<tr>
<td>Bronchoscopically (B) obtained specimens</td>
<td></td>
</tr>
<tr>
<td>Bronchoalveolar lavage (B-BAL)</td>
<td>≥10⁴ cfu/ml</td>
</tr>
<tr>
<td>Protected BAL (B-PBAL)</td>
<td>≥10⁴ cfu/ml</td>
</tr>
<tr>
<td>Protected specimen brushing (B-PSB)</td>
<td>≥10³ cfu/ml</td>
</tr>
<tr>
<td>Nonbronchoscopically (NB) obtained</td>
<td></td>
</tr>
<tr>
<td>specimens</td>
<td></td>
</tr>
<tr>
<td>NB-BAL</td>
<td>&gt;10⁷ cfu/ml</td>
</tr>
<tr>
<td>NB-PSB</td>
<td>≥10³ cfu/ml</td>
</tr>
</tbody>
</table>

cfu = colony forming units  
g = gram  
ml = milliliter

**COMMENT:**  
* Open-lung biopsy specimens and immediate post-mortem specimens obtained by transthoracic or transbronchial biopsy

**Numerator Data:** The *Pneumonia (PNEU)* from (CDC 57.111) is used to collect and report each VAP that is identified during the month selected for surveillance. The *Instructions for Completion of Pneumonia Form* (Tables of Instructions, Tables 4 and 2a) includes brief instructions for collection and entry of each data element on the form. The pneumonia form includes patient demographic information and information on whether or not mechanically assisted ventilation was present. Additional data include the specific criteria met for identifying pneumonia, whether the patient developed a secondary bloodstream infection, whether the patient died, and the organisms isolated from cultures and their antimicrobial susceptibilities.

**Denominator data:** Device days and patient days are used for denominators (see **Chapter 16** Key Terms). Ventilator days, which are the number of patients managed with a ventilatory device, are collected daily, at the same time each day, according to the chosen location using the appropriate form (CDC 57.116, 57.117, and 57.118). These daily counts are summed and only the total for the month is entered into NHSN. Ventilator and patient days are collected for each of the locations monitored. When denominator data are available from electronic sources (e.g., ventilator days from respiratory therapy),
these sources may be used as long as the counts are not substantially different (+/- 5%) from manually collected counts.

Data Analyses: The SIR is calculated by dividing the number of observed infections by the number of expected infections. The number of expected infections, in the context of statistical prediction, is calculated using PNEU rates from a standard population during a baseline time period as reported in the NHSN Report.

NOTE: The SIR will be calculated only if the number of expected HAIs (numExp) is ≥ 1.

\[
\text{SIR} = \frac{\text{Observed (O) HAIs}}{\text{Expected (E) HAIs}}
\]

While the PNEU SIR can be calculated for single locations, the measure also allows you to summarize your data by multiple locations, adjusting for differences in the incidence of infection among the location types. For example, you will be able to obtain one PNEU SIR adjusting for all locations reported. Similarly, you can obtain one PNEU SIR for all specialty care areas in your facility.

The VAP rate per 1000 ventilator days is calculated by dividing the number of VAPs by the number of ventilator days and multiplying the result by 1000. The Ventilator Utilization Ratio is calculated by dividing the number of ventilator days by the number of patient days. These calculations will be performed separately for the different types of ICUs, SCAs, and other locations in the institution, as well as by each birthweight category in NICUs.


### SSI

<table>
<thead>
<tr>
<th>event category</th>
<th>full term</th>
<th>short term</th>
<th>specifications</th>
<th>standard</th>
</tr>
</thead>
<tbody>
<tr>
<td>healthcare-associated infection</td>
<td>surgical site infection</td>
<td>SSI</td>
<td>Results in serious physical injury, death or higher level of care including but not limited to prolonged hospital stay, IV antibiotics</td>
<td>AHRQ, NHSN</td>
</tr>
</tbody>
</table>

### Implementation guidance

There will be surgical site infections that are reported to NHSN but are NOT sentinel events.

NHSN guidance included on the next few pages.

### Examples

*not a sentinel event*: A 41 year-old male presents to hospital to have an ankle fusion. The fusion is done successfully. At day 2 prior to discharge, the wound is noted to be red, painful, and warm to the touch. There is neither purulence nor fluctuation. The patient is treated with post-operative Keflex and discharged. Patient follow up with the surgeon reveals a well-healed wound. This would meet the definition of a superficial, incisional SSI but not a sentinel event.
**Surgical Site Infection (SSI) Event**

**Introduction:** In 2002, in the United States, an estimated 14 million NHSN operative procedures were performed (CDC unpublished data). SSIs were the second most common healthcare-associated infection, accounting for 17% of all HAIs among hospitalized patients. A similar rate was obtained from NHSN hospitals reporting data in 2006-2008 (15,862 SSI following 830,748 operative procedures) (CDC, unpublished data) with an overall rate of nearly 2%.

While advances have been made in infection control practices, including improved operating room ventilation, sterilization methods, barriers, surgical technique, and availability of antimicrobial prophylaxis, SSIs remain a substantial cause of morbidity and mortality among hospitalized patients. In one study, among nearly 100,000 HAIs reported in one year, deaths were associated with SSIs in more than 8,000 cases.

Surveillance of SSI with feedback of appropriate data to surgeons has been shown to be an important component of strategies to reduce SSI risk. A successful surveillance program includes the use of epidemiologically-sound infection definitions and effective surveillance methods, stratification of SSI rates according to risk factors associated with SSI development, and data feedback. Recommendations are outlined in the CDC’s *Guideline for Prevention of Surgical Site Infection, 1999.*

**Settings:** Surveillance will occur with surgical patients in any inpatient/outpatient setting where the selected NHSN operative procedure(s) are performed.

**Requirements:** Select at least one NHSN operative procedure category (Table 1) and indicate this on the *Patient Safety Monthly Reporting Plan* (CDC 57.106). Collect numerator and denominator data on all selected procedure categories for at least one month.

The *International Classification of Diseases, 9th Revision Clinical Modifications* (ICD-9-CM) codes, which are defined by the ICD-9 Coordination and Maintenance Committee of the National Center for Health Statistics and the Centers for Medicare and Medicaid Services (CMS), are developed as a tool for classification of morbidity data. The preciseness of the data, as well as their wide use, allows their use in grouping surgery types for the purpose of determining SSI rates. ICD-9-CM codes are updated annually in October and NHSN operative procedure categories are subsequently updated and changes shared with NHSN users. Table 1: NHSN Operative Procedure Category Mappings to ICD-9-CM Codes, below, outlines operative procedures and their grouping into NHSN operative procedure categories according to ICD-9-CM codes. A brief description of the types of operations contained in the NHSN operative procedure categories is also provided.
Table 1. NHSN Operative Procedure Category Mappings to ICD-9-CM Codes

<table>
<thead>
<tr>
<th>Legacy Code</th>
<th>Operative Procedure</th>
<th>Description</th>
<th>ICD-9-CM Codes</th>
</tr>
</thead>
<tbody>
<tr>
<td>AAA</td>
<td>Abdominal aortic aneurysm repair</td>
<td>Resection of abdominal aorta with anastomosis or replacement</td>
<td>38.34, 38.44, 38.64</td>
</tr>
<tr>
<td>AMP</td>
<td>Limb amputation</td>
<td>Total or partial amputation or disarticulation of the upper or lower limbs, including digits</td>
<td>84.00-84.19, 84.91</td>
</tr>
<tr>
<td>APPY</td>
<td>Appendix surgery</td>
<td>Operation of appendix (not incidental to another procedure)</td>
<td>47.01, 47.09, 47.2, 47.91, 47.92, 47.99</td>
</tr>
<tr>
<td>AVSD</td>
<td>Shunt for dialysis</td>
<td>Arteriovenostomy for renal dialysis</td>
<td>39.27, 39.42</td>
</tr>
<tr>
<td>BILI</td>
<td>Bile duct, liver or pancreatic surgery</td>
<td>Excision of bile ducts or operative procedures on the biliary tract, liver or pancreas (does not include operations only on gallbladder)</td>
<td>50.0, 50.12, 50.14, 50.21-50.23, 50.25, 50.26, 50.29, 50.3, 50.4, 50.61, 50.69, 51.31-51.37, 51.39, 51.41-51.43, 51.49, 51.51, 51.59, 51.61-51.63, 51.69, 51.71, 51.72, 51.79, 51.81-51.83, 51.89, 51.91-51.95, 51.99, 52.09, 52.12, 52.22, 52.3, 52.4, 52.51-52.53, 52.59-52.6, 52.7, 52.92, 52.95, 52.96, 52.99</td>
</tr>
<tr>
<td>BRST</td>
<td>Breast surgery</td>
<td>Excision of lesion or tissue of breast including radical, modified, or quadrant resection, lumpectomy, incisional biopsy, or mammoplasty.</td>
<td>85.12, 85.20-85.23, 85.31-85.36, 85.41-85.48, 85.50, 85.53-85.55, 85.6, 85.70-85.76, 85.79, 85.93-85.96</td>
</tr>
<tr>
<td>CARD</td>
<td>Cardiac surgery</td>
<td>Procedures on the valves or septum of heart; does not include coronary artery bypass graft, surgery on vessels, heart transplantation, or pacemaker implantation</td>
<td>35.00-35.04, 35.10-35.14, 35.20-35.28, 35.31-35.35, 35.39, 35.42, 35.50, 35.51, 35.53, 35.54, 35.60-35.63, 35.70-35.73, 35.81-35.84, 35.91-35.95, 35.98-35.99, 37.10-37.12, 37.31-37.33, 37.35-37.37, 37.41, 37.49, 37.60*</td>
</tr>
<tr>
<td>CEA</td>
<td>Carotid endarterectomy</td>
<td>Endarterectomy on vessels of head and neck (includes carotid artery and jugular vein)</td>
<td>38.12</td>
</tr>
</tbody>
</table>
## Legacy Code | Operative Procedure | Description | ICD-9-CM Codes
--- | --- | --- | ---
CBGB | Coronary artery bypass graft with **both** chest and donor site incisions | Chest procedure to perform direct revascularization of the heart; includes obtaining suitable vein from donor site for grafting. | 36.10-36.14, 36.19
CBGC | Coronary artery bypass graft with chest incision only | Chest procedure to perform direct vascularization of the heart using, for example the internal mammary (thoracic) artery | 36.15-36.17, 36.2
CHOL | Gallbladder surgery | Cholecystectomy and cholecystotomy | 51.03, 51.04, 51.13, 51.21-51.24
COLO | Colon surgery | Incision, resection, or anastomosis of the large intestine; includes large-to-small and small-to-large bowel anastomosis; does not include rectal operations | 17.31-17.36, 17.39, 45.03, 45.26, 45.41, 45.49, 45.52, 45.71-45.76, 45.79, 45.81-45.83, 45.92-45.95, 46.03, 46.04, 46.10, 46.11, 46.13, 46.14, 46.43, 46.52, 46.75, 46.76, 46.94
CRAN | Craniotomy | Excision repair, or exploration of the brain or meninges; does not include taps or punctures | 01.12, 01.14, 01.20-01.25, 01.28, 01.29, 01.31, 01.32, 01.39, 01.41, 01.42, 01.51-01.53, 01.59, 02.11-02.14, 02.91-02.93, 07.51-07.54, 07.59, 07.61-07.65, 07.68, 07.69, 07.71, 07.72, 07.79, 38.01, 38.11, 38.31, 38.41, 38.51, 38.61, 38.81, 39.28
CSEC | Cesarean section | Obstetrical delivery by Cesarean section | 74.0, 74.1, 74.2, 74.4, 74.91, 74.99
FUSN | Spinal fusion | Immobilization of spinal column | 81.00-81.08
FX | Open reduction of fracture | Open reduction of fracture or dislocation of long bones with or without internal or external fixation; does not include placement of joint prosthesis | 79.21, 79.22, 79.25, 79.26, 79.31, 79.32, 79.35, 79.36, 79.51, 79.52, 79.55, 79.56
GAST | Gastric surgery | Incision or excision of stomach; includes subtotal or total gastrectomy; does not include vagotomy and fundoplication | 43.0, 43.42, 43.49, 43.5, 43.6, 43.7, 43.81, 43.89, 43.91, 43.99, 44.15, 44.21, 44.29, 44.31, 44.38-44.42, 44.49, 44.5, 44.61-44.65, 44.68-44.69, 44.95-44.98
<table>
<thead>
<tr>
<th>Legacy Code</th>
<th>Operative Procedure</th>
<th>Description</th>
<th>ICD-9-CM Codes</th>
</tr>
</thead>
<tbody>
<tr>
<td>HER</td>
<td>Herniorrhaphy</td>
<td>Repair of inguinal, femoral, umbilical, or anterior abdominal wall hernia; does not include repair of diaphragmatic or hiatal hernia or hernias at other body sites.</td>
<td>17.11-17.13, 17.21-17.24, 53.00-53.05, 53.10-53.17, 53.21, 53.29, 53.31, 53.39, 53.41-53.43, 53.49, 53.51, 53.59, 53.61-53.63, 53.69</td>
</tr>
<tr>
<td>HPRO</td>
<td>Hip prosthesis</td>
<td>Arthroplasty of hip</td>
<td>00.70-00.73, 00.85-00.87, 81.51-81.53</td>
</tr>
<tr>
<td>HTP</td>
<td>Heart transplant</td>
<td>Transplantation of heart</td>
<td>37.51-37.55</td>
</tr>
<tr>
<td>HYST</td>
<td>Abdominal hysterectomy</td>
<td>Abdominal approach with uterine removal</td>
<td>68.31, 68.39, 68.41, 68.49, 68.61, 68.69</td>
</tr>
<tr>
<td>KPRO</td>
<td>Knee prosthesis</td>
<td>Arthroplasty of knee</td>
<td>00.80-00.84, 81.54, 81.55</td>
</tr>
<tr>
<td>KTP</td>
<td>Kidney transplant</td>
<td>Transplantation of kidney</td>
<td>55.61, 55.69</td>
</tr>
<tr>
<td>LAM</td>
<td>Laminectomy</td>
<td>Exploration or decompression of spinal cord through excision or incision into vertebral structures</td>
<td>03.01, 03.02, 03.09, 80.50, 80.51, 80.53, 80.54†, 80.59, 84.60-84.69, 84.80-84.85</td>
</tr>
<tr>
<td>LTP</td>
<td>Liver transplant</td>
<td>Transplantation of liver</td>
<td>50.51, 50.59</td>
</tr>
<tr>
<td>NECK</td>
<td>Neck surgery</td>
<td>Major excision or incision of the larynx and radical neck dissection; does not include thyroid and parathyroid operations.</td>
<td>30.1, 30.21, 30.22, 30.29, 30.3, 30.4, 31.45, 40.40-40.42</td>
</tr>
<tr>
<td>NEPH</td>
<td>Kidney surgery</td>
<td>Resection or manipulation of the kidney with or without removal of related structures</td>
<td>55.01, 55.02, 55.11, 55.12, 55.24, 55.31, 55.32, 55.34, 55.35, 55.39, 55.4, 55.51, 55.52, 55.54, 55.91</td>
</tr>
<tr>
<td>PACE</td>
<td>Pacemaker surgery</td>
<td>Insertion, manipulation or replacement of pacemaker</td>
<td>00.50-00.54, 17.51, 17.52, 37.70-37.77, 37.79-37.83, 37.85-37.87, 37.89, 37.94-37.99</td>
</tr>
<tr>
<td>PRST</td>
<td>Prostate surgery</td>
<td>Suprapubic, retropubic, radical, or perineal excision of the prostate; does not include transurethral</td>
<td>60.12, 60.3, 60.4, 60.5, 60.61, 60.62, 60.69</td>
</tr>
<tr>
<td>Legacy Code</td>
<td>Operative Procedure</td>
<td>Description</td>
<td>ICD-9-CM Codes</td>
</tr>
<tr>
<td>-------------</td>
<td>---------------------</td>
<td>------------------------------------------------------------------------------</td>
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</tr>
<tr>
<td>PVBY</td>
<td>Peripheral vascular bypass surgery</td>
<td>Bypass operations on peripheral arteries</td>
<td>39.29</td>
</tr>
<tr>
<td>RFUSN</td>
<td>Refusion of spine</td>
<td>Refusion of spine</td>
<td>81.30-81.39</td>
</tr>
<tr>
<td>SB</td>
<td>Small bowel surgery</td>
<td>Incision or resection of the small intestine; does not include small-to-large bowel anastomosis</td>
<td>45.01, 45.02, 45.15, 45.31-45.34, 45.51, 45.61-45.63, 45.91, 46.01, 46.02, 46.20-46.24, 46.31, 46.39, 46.41, 46.51, 46.71-46.74, 46.93</td>
</tr>
<tr>
<td>SLE</td>
<td>Spleen surgery</td>
<td>Resection or manipulation of spleen</td>
<td>41.2, 41.33, 41.41-41.43, 41.5, 41.93, 41.95, 41.99</td>
</tr>
<tr>
<td>THOR</td>
<td>Thoracic surgery</td>
<td>Noncardiac, nonvascular thoracic surgery; includes pneumonectomy and hiatal hernia repair or diaphragmatic hernia repair (except through abdominal approach.)</td>
<td>32.09, 32.1, 32.20-32.23, 32.25, 32.26, 32.29, 32.30, 32.39, 32.41, 32.49, 32.50, 32.59, 32.6, 32.9, 33.0, 33.1, 33.20, 33.25, 33.28, 33.31-33.34, 33.39, 33.41-33.43, 33.48, 33.49, 33.98, 33.99, 34.01-34.03, 34.06, 34.1, 34.20, 34.26, 34.3, 34.4, 34.51, 34.52, 34.59, 34.6, 34.81-34.84, 34.89, 34.93, 34.99, 53.80-53.84</td>
</tr>
<tr>
<td>THYR</td>
<td>Thyroid and/or parathyroid surgery</td>
<td>Resection or manipulation of thyroid and/or parathyroid</td>
<td>06.02, 06.09, 06.12, 06.2, 06.31, 06.39, 06.4, 06.50-06.52, 06.6, 06.7, 06.81, 06.89, 06.91-06.95, 06.98, 06.99</td>
</tr>
<tr>
<td>VHYS</td>
<td>Vaginal hysterectomy</td>
<td>Vaginal approach with uterine removal</td>
<td>68.51, 68.59, 68.71, 68.79</td>
</tr>
<tr>
<td>VSHN</td>
<td>Ventricular shunt</td>
<td>Ventricular shunt operations, including revision and removal of shunt</td>
<td>02.2, 02.31-02.35, 02.39, 02.42, 02.43, 54.95*</td>
</tr>
<tr>
<td>XLAP</td>
<td>Abdominal surgery</td>
<td>Abdominal operations not involving the gastrointestinal tract or biliary system includes diaphragmatic hernia repair through abdominal approach.</td>
<td>53.71, 53.72, 53.75, 54.0, 54.11, 54.12, 54.19, 54.3, 54.4, 54.51, 54.59, 54.61, 54.63, 54.64, 54.71-54.75, 54.92, 54.93</td>
</tr>
</tbody>
</table>
*NOTE: The procedure represented by this ICD-9-CM code can be performed in a number of ways. However, as for all surgeries, if, at the end of the procedure, the skin incision edges do not meet because of wires, devices or other objects extruding through the incision, the incision is not considered primarily closed. Therefore the procedure is not considered an NHSN operative procedure and any subsequent infection is not considered a procedure-associated infection (i.e., not an SSI or PPP).

†NOTE: If this procedure is performed percutaneously, it is not considered an NHSN operative procedure and should not be included in LAM denominator data.

^ NOTE: Include only if this procedure involves ventricular shunt.

For a complete mapping of all ICD-9-CM codes to their assignment as an NHSN operative procedure category, a surgical procedure other than an NHSN operative procedure (OTH), or a non-operative procedure (NO), see ICD-9-CM Procedure Code Mapping to NHSN Operative Procedure Categories at [http://www.cdc.gov/nhsn/library.html](http://www.cdc.gov/nhsn/library.html).

Definitions:

An **NHSN operative procedure** is a procedure

1) that is performed on a patient who is an NHSN inpatient or an NHSN outpatient; 2) takes place during an operation (defined as a single trip to the operating room (OR) where a surgeon makes at least one incision through the skin or mucous membrane, including laparoscopic approach, and closes the incision before the patient leaves the OR; and 3) that is included in Table 1.

*NOTE: If the skin incision edges do not meet because of wires or devices or other objects extruding through the incision, the incision is not considered primarily closed and therefore the procedure is not considered an operation. Further, any subsequent infection is not considered a procedure-associated infection (i.e., not an SSI or PPP).

**NHSN Inpatient:** A patient whose date of admission to the healthcare facility and the date of discharge are different calendar days.

**NHSN Outpatient:** A patient whose date of admission to the healthcare facility and date of discharge are the same calendar day.

**Operating Room (OR):** A patient care area that met the Facilities Guidelines Institute’s (FGI) or American Institute of Architects’ (AIA) criteria for an operating room when it was constructed or renovated. This may include an operating room, C-Section room, interventional radiology room, or a cardiac catheterization lab.

**Implant:** A nonhuman-derived object, material, or tissue that is permanently placed in a patient during an operative procedure and is not routinely manipulated for diagnostic or therapeutic purposes. Examples include: porcine or synthetic heart valves, mechanical heart, metal rods, mesh, sternal wires, screws, cements, internal staples, hemoclips, and other devices. Non-absorbable
sutures are excluded because Infection Preventionists may not easily identify and/or differentiate the soluble nature of suture material used.

Transplant: Human cells, tissues, organs, or cellular- or tissue-based products that are placed into a human recipient via grafting, infusion, or transfer. Examples include: heart valves, organs, ligaments, bone, blood vessels, skin, corneas, and bone marrow cells. Autologous or “autograft” transplants are products that originate from the patient’s own body. Non-autologous or “allograft” transplants are tissues or other products derived from another human body, either a donor cadaver or a live donor.

REPORTING INSTRUCTIONS:
- Some products are a combination of human- and nonhuman-derived materials, such as demineralized human bone matrix with porcine gel carrier. When placed in a patient during an operative procedure, indicate “Yes” for both the Implant and Non-autologous Transplant fields.
- Some operative procedures involve placement of both autologous and non-autologous products. For these procedures, indicate “Yes” for Non-autologous Transplant field.

A superficial incisional SSI must meet one of the following criteria:
Infection occurs within 30 days after the operative procedure and involves only skin and subcutaneous tissue of the incision and patient has at least one of the following:
  a. purulent drainage from the superficial incision.
  b. organisms isolated from an aseptically obtained culture of fluid or tissue from the superficial incision.
  c. at least one of the following signs or symptoms of infection: pain or tenderness, localized swelling, redness, or heat, and superficial incision is deliberately opened by surgeon, and is culture-positive or not cultured. A culture-negative finding does not meet this criterion.
  d. diagnosis of superficial incisional SSI by the surgeon or attending physician.

NOTE: There are two specific types of superficial incisional SSIs:
1. Superficial Incisional Primary (SIP) – a superficial incisional SSI that is identified in the primary incision in a patient that has had an operation with one or more incisions (e.g., C-section incision or chest incision for CBGB)
2. Superficial Incisional Secondary (SIS) – a superficial incisional SSI that is identified in the secondary incision in a patient that has had an operation with more than one incision (e.g., donor site [leg] incision for CBGB)

REPORTING INSTRUCTIONS:
- Do not report a stitch abscess (minimal inflammation and discharge confined to the points of suture penetration) as an infection.
- Do not report a localized stab wound infection as SSI. While it would be considered either a skin (SKIN) or soft tissue (ST) infection, depending on its depth, it is not reportable under this module.
- “Cellulitis”, by itself, does not meet the criteria for Superficial Incisional SSI.
- If the incisional site infection involves or extends into the fascial and muscle layers, report as a deep-incisional SSI.
- Classify infection that involves both superficial and deep incision sites as deep incisional SSI.
- An infected circumcision site in newborns is classified as CIRC. Circumcision is not an NHSN operative procedure. CIRC is not reportable under this module.
- An infected burn wound is classified as BURN and is not reportable under this module.

A **deep incisional SSI** must meet one of the following criteria:

Infection occurs within 30 days after the operative procedure if no implant is left in place or within one year if implant is in place and the infection appears to be related to the operative procedure and involves deep soft tissues (e.g., fascial and muscle layers) of the incision and patient has at least one of the following:

- a. purulent drainage from the deep incision but not from the organ/space component of the surgical site
- b. a deep incision spontaneously dehisces or is deliberately opened by a surgeon and is culture-positive or not cultured and the patient has at least one of the following signs or symptoms: fever (>38°C), or localized pain or tenderness. A culture-negative finding does not meet this criterion.
- c. an abscess or other evidence of infection involving the deep incision is found on direct examination, during reoperation, or by histopathologic or radiologic examination
- d. diagnosis of a deep incisional SSI by a surgeon or attending physician.

NOTE: There are two specific types of deep incisional SSIs:

1. **Deep Incisional Primary (DIP)** – a deep incisional SSI that is identified in a primary incision in a patient that has had an operation with one or more incisions (e.g., C-section incision or chest incision for CBGB)

2. **Deep Incisional Secondary (DIS)** – a deep incisional SSI that is identified in the secondary incision in a patient that has had an operation with more than one incision (e.g., donor site [leg] incision for CBGB)

**REPORTING INSTRUCTIONS:**

- Classify infection that involves both superficial and deep incision sites as deep incisional SSI.

An **organ/space SSI** involves any part of the body, excluding the skin incision, fascia, or muscle layers, that is opened or manipulated during the operative procedure. Specific sites are assigned to organ/space SSI to further identify the location of the infection. The table below lists the specific sites that must be used to differentiate organ/space SSI. An example is appendectomy with
subsequent subdiaphragmatic abscess, which would be reported as an organ/space SSI at the intraabdominal specific site (SSI-IAB). Specific sites of organ/space (Table 2) have specific criteria which must be met in order to qualify as an NHSN event. These criteria are in addition to the general criteria for organ/space SSI and can be found in Chapter 17.

An organ/space SSI must meet one of the following criteria:
Infection occurs within 30 days after the operative procedure if no implant is left in place or within one year if implant is in place and the infection appears to be related to the operative procedure and infection involves any part of the body, excluding the skin incision, fascia, or muscle layers, that is opened or manipulated during the operative procedure and patient has at least one of the following:
   a. purulent drainage from a drain that is placed through a stab wound into the organ/space
   b. organisms isolated from an aseptically obtained culture of fluid or tissue in the organ/space
   c. an abscess or other evidence of infection involving the organ/space that is found on direct examination, during reoperation, or by histopathologic or radiologic examination
   d. diagnosis of an organ/space SSI by a surgeon or attending physician.

REPORTING INSTRUCTIONS:
- Occasionally an organ/space infection drains through the incision. Such infection generally does not involve reoperation and is considered a complication of the incision. Therefore, classify it as a deep incisional SSI.
- Report mediastinitis following cardiac surgery that is accompanied by osteomyelitis as SSI-MED rather than SSI-BONE.
- If meningitis (MEN) and a brain abscess (IC) are present together after operation, report as SSI-IC.
- Report CSF shunt infection as SSI-MEN if it occurs ≤ 1 year of placement; if later or after manipulation/access, it is considered CNS-MEN and is not reportable under this manual.
- Report spinal abscess with meningitis as SSI-MEN following spinal surgery.
- Episiotomy is not considered an operative procedure in NHSN.

Table 2. Specific sites of an organ/space SSI. Criteria for these sites can be found in the NHSN Help System (must be logged in to NHSN) or Chapter 17.

<table>
<thead>
<tr>
<th>Code</th>
<th>Site</th>
<th>Code</th>
<th>Site</th>
</tr>
</thead>
<tbody>
<tr>
<td>BONE</td>
<td>Osteomyelitis</td>
<td>JNT</td>
<td>Joint or bursa</td>
</tr>
<tr>
<td>BRST</td>
<td>Breast abscess or mastitis</td>
<td>LUNG</td>
<td>Other infections of the respiratory tract</td>
</tr>
<tr>
<td>CARD</td>
<td>Myocarditis or pericarditis</td>
<td>MED</td>
<td>Mediastinitis</td>
</tr>
<tr>
<td>DISC</td>
<td>Disc space</td>
<td>MEN</td>
<td>Meningitis or ventriculitis</td>
</tr>
<tr>
<td>EAR</td>
<td>Ear, mastoid</td>
<td>ORAL</td>
<td>Oral cavity (mouth, tongue, or gums)</td>
</tr>
<tr>
<td>EMET</td>
<td>Endometritis</td>
<td>OREP</td>
<td>Other infections of the male or female reproductive tract</td>
</tr>
</tbody>
</table>
**Procedures-associated Events**

**SSI**

<table>
<thead>
<tr>
<th>Code</th>
<th>Site</th>
<th>Code</th>
<th>Site</th>
</tr>
</thead>
<tbody>
<tr>
<td>ENDO</td>
<td>Endocarditis</td>
<td>OUTI</td>
<td>Other infections of the urinary tract</td>
</tr>
<tr>
<td>EYE</td>
<td>Eye, other than conjunctivitis</td>
<td>SA</td>
<td>Spinal abscess without meningitis</td>
</tr>
<tr>
<td>GIT</td>
<td>GI tract</td>
<td>SINU</td>
<td>Sinusitis</td>
</tr>
<tr>
<td>HEP</td>
<td>Hepatitis</td>
<td>UR</td>
<td>Upper respiratory tract</td>
</tr>
<tr>
<td>IAB</td>
<td>Intraabdominal, not specified else-where</td>
<td>VASC</td>
<td>Arterial or venous infection</td>
</tr>
<tr>
<td>IC</td>
<td>Intracranial, brain abscess or dura</td>
<td>VCUF</td>
<td>Vaginal cuff</td>
</tr>
</tbody>
</table>

**Numerator Data:** All patients having the selected operative procedure are monitored for signs of SSI. The *Surgical Site Infection (SSI)* form (CDC 57.120) is completed for each such patient found to have an SSI.

**NOTES:**

1. If a patient has several NHSN operative procedures prior to an infection, report the operative procedure code of the operation that was performed most closely in time prior to the infection date, unless there is evidence that the infection is associated with a different operation.

2. If a procedure from more than one NHSN operative procedure category was done through a single incision, attempt to determine the procedure that is thought to be associated with the infection. If it is not clear (as is often the case when the infection is a superficial incisional SSI), or if the infection site being reported is not an SSI, use the NHSN Principal Operative Procedure Category Selection Lists (Table 3) to select which operative procedure to report.

**Table 3. NHSN Principal Operative Procedure Category Selection Lists**

<table>
<thead>
<tr>
<th>Priority</th>
<th>Code</th>
<th>Abdominal Operations</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>SB</td>
<td>Small bowel surgery</td>
</tr>
<tr>
<td>2</td>
<td>KTP</td>
<td>Kidney transplant</td>
</tr>
<tr>
<td>3</td>
<td>LTP</td>
<td>Liver transplant</td>
</tr>
<tr>
<td>4</td>
<td>BILI</td>
<td>Bile duct, liver or pancreatic surgery</td>
</tr>
<tr>
<td>5</td>
<td>REC</td>
<td>Rectal surgery</td>
</tr>
<tr>
<td>6</td>
<td>COLO</td>
<td>Colon surgery</td>
</tr>
<tr>
<td>7</td>
<td>GAST</td>
<td>Gastric surgery</td>
</tr>
<tr>
<td>8</td>
<td>CSEC</td>
<td>Cesarean section</td>
</tr>
<tr>
<td>9</td>
<td>SPLE</td>
<td>Spleen surgery</td>
</tr>
<tr>
<td>10</td>
<td>APPY</td>
<td>Appendix surgery</td>
</tr>
<tr>
<td>11</td>
<td>HYST</td>
<td>Abdominal hysterectomy</td>
</tr>
<tr>
<td>12</td>
<td>VHYS</td>
<td>Vaginal Hysterectomy</td>
</tr>
<tr>
<td>13</td>
<td>OVRY</td>
<td>Ovarian surgery</td>
</tr>
<tr>
<td>14</td>
<td>HER</td>
<td>Herniorrhaphy</td>
</tr>
</tbody>
</table>

The following lists are derived from Table 1, NHSN Operative Procedure Categories. The operative procedures with the highest risk of surgical site infection are listed before those with a lower risk.
The following lists are derived from Table 1, NHSN Operative Procedure Categories. The operative procedures with the highest risk of surgical site infection are listed before those with a lower risk.

<table>
<thead>
<tr>
<th>Priority</th>
<th>Code</th>
<th>Procedure</th>
</tr>
</thead>
<tbody>
<tr>
<td>15</td>
<td>CHOL</td>
<td>Gall bladder surgery</td>
</tr>
<tr>
<td>16</td>
<td>AAA</td>
<td>Abdominal aortic aneurysm repair</td>
</tr>
<tr>
<td>17</td>
<td>NEPH</td>
<td>Kidney surgery</td>
</tr>
<tr>
<td>18</td>
<td>XLAP</td>
<td>Laparotomy</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Priority</th>
<th>Code</th>
<th>Thoracic Operations</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>HTP</td>
<td>Heart transplant</td>
</tr>
<tr>
<td>2</td>
<td>CBGB</td>
<td>Coronary artery bypass graft with donor incision(s)</td>
</tr>
<tr>
<td>3</td>
<td>CBGC</td>
<td>Coronary artery bypass graft, chest incision only</td>
</tr>
<tr>
<td>4</td>
<td>CARD</td>
<td>Cardiac surgery</td>
</tr>
<tr>
<td>5</td>
<td>THOR</td>
<td>Thoracic surgery</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Priority</th>
<th>Code</th>
<th>Neurosurgical (Spine) Operations</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>RFUSN</td>
<td>Refusion of spine</td>
</tr>
<tr>
<td>2</td>
<td>FUSN</td>
<td>Spinal fusion</td>
</tr>
<tr>
<td>3</td>
<td>LAM</td>
<td>Laminectomy</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Priority</th>
<th>Code</th>
<th>Neurosurgical (Brain) Operations</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>VSHN</td>
<td>Ventricular shunt</td>
</tr>
<tr>
<td>2</td>
<td>CRAN</td>
<td>Craniotomy</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Priority</th>
<th>Code</th>
<th>Neck Operations</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>NECK</td>
<td>Neck surgery</td>
</tr>
<tr>
<td>2</td>
<td>THYR</td>
<td>Thyroid and or parathyroid surgery</td>
</tr>
</tbody>
</table>

The Instructions for Completion of Surgical Site Infection form (Tables of Instructions, Tables 12 and 2a) includes brief instructions for collection and entry of each data element on the form. The SSI form includes patient demographic information and information about the operative procedure, including the date and type of procedure. Information about the SSI includes the date of SSI, specific criteria met for identifying the SSI, when the SSI was detected, whether the patient developed a secondary bloodstream infection, whether the patient died, and the organisms isolated from cultures and the organisms’ antimicrobial susceptibilities.

**Denominator Data:** For all patients having a procedure selected for surveillance during the month, complete the Denominator for Procedure form (CDC 57.121). The data are collected individually for each operative procedure performed during the month specified on the Patient Safety Monthly Surveillance Plan (CDC 57.106). The Instructions for Completion of Denominator for Procedure form (Tables of Instructions, Table 13) includes brief instructions for collection and entry of each data element on the form.

NOTES:
1. If procedures in more than one NHSN operative procedure category are performed during the same trip to the OR even if performed through the same incision, a Denominator for Procedure (CDC 57.121) record is reported for each operative procedure being monitored. For example, if a CARD and CBGC are done through the same incision, a Denominator for Procedure record is reported for each.

2. EXCEPTION: If a patient has both a CBGC and CBGB during the same trip to the OR, report only as a CBGB. Only report as a CBGC when there is a chest incision only. CBGB and CBGC are never reported for the same patient for the same trip to the OR. For bilateral operative procedures see #4 below.

3. If procedures of different ICD-9-CM codes from the same NHSN Operative Procedure Category are performed through the same incision, record only one procedure for that category. For example, if your facility is performing surveillance for both CBGB and CARD procedures, and a patient undergoes an aortocoronary bypass of one coronary vessel (36.11, CBGB) and the replacement of both the mitral and tricuspid valves (35.23 and 35.27, both CARD) during the same trip to the OR, you would complete a Denominator for Procedure record for the CBGB and another for the CARD.

4. If more than one NHSN operative procedure category is performed through the same incision, record the combined duration of all procedures, which is the time from skin incision to primary closure.

5. For bilateral operative procedures (e.g., KPRO), two separate Denominator for Procedure (CDC 57.121) records are completed. To document the duration of the procedure, indicate the incision time to closure time for each procedure separately or, alternatively, take the total time for both procedures and split it evenly between the two. See “5” below.

6. Laparoscopic hernia repairs are considered one procedure, regardless of the number of hernias that are repaired in that trip to the OR. In most cases there will be only one incision time documented for this procedure. If more than one time is documented, total the durations. In this situation, if more than one of the incisions should become infected, only report as a single SSI. Open [i.e., non-laparoscopic] hernia repairs are reported as one procedure for each hernia repaired via a separate incision, i.e., if two incisions are made to repair two defects, then two procedures will be reported. It is anticipated that separate incision times will be recorded for these procedures. If not, take the total time for both procedures and split it evenly between the two.

7. If a patient goes to the OR more than once during the same admission and another procedure is performed through the same incision within 24 hours of the original operative incision, report only one procedure on the Denominator for Procedure (CDC 57.121) form combining the durations for both procedures. For example, a patient has a CBGB lasting 4 hours. He returns to the OR six hours later to correct a bleeding vessel. The surgeon reopens the initial incision, makes the repairs, and recloses in 1.5 hours. Record the operative procedure as one CBGB and the duration of operation as 5 hour 30 minutes. If the wound class has changed, report the higher wound class. If the ASA class has changed, report the higher ASA class.
**Data Analyses:** The SIR is calculated by dividing the number of observed infections by the number of expected infections. The number of expected infections, in the context of statistical prediction, is calculated using SSI probabilities estimated from multivariate logistic regression models constructed from NHSN data during a baseline time period to represent a standard population.

**NOTE:** The SIR will be calculated only if the number of expected HAIs (numExp) is ≥ 1.

\[
SIR = \frac{\text{Observed (O) HAIs}}{\text{Expected (E) HAIs}}
\]

While the SSI SIR can be calculated for single procedure categories, and for specific surgeons, the measure also allows you to summarize your data across multiple procedure categories, while adjusting for differences in the estimated probability of infection among the patients included across the procedure categories. For example, you will be able to obtain one SSI SIR adjusting for all procedures reported. Alternatively, you can obtain one SSI SIR for all colon surgeries (COLO) only within your facility.

SSI rates per 100 operative procedures are calculated by dividing the number of SSIs by the number of specific operative procedures and multiplying the results by 100. SSI will be included in the numerator of a rate based on the date of procedure, not the date of event. Rate calculations can be performed separately for the different types of operative procedures and stratified by the basic risk index. SSI rate calculation options are available in the advanced analysis feature of the NHSN application.

- **Basic SSI Risk Index.** The index used in NHSN assigns surgical patients into categories based on the presence of three major risk factors:

  1. Operation lasting more than the duration cut point hours, where the duration cut point is the approximate 75th percentile of the duration of surgery in minutes for the operative procedure.

  2. Contaminated (Class 3) or Dirty/infected (Class 4) wound class.

  3. ASA classification of 3, 4, or 5.

  The patient’s SSI risk category is simply the number of these factors present at the time of the operation.

---


## CAUTI

<table>
<thead>
<tr>
<th>Event Category</th>
<th>Full Term</th>
<th>Short Term</th>
<th>Specifications</th>
<th>Standard</th>
</tr>
</thead>
<tbody>
<tr>
<td>Healthcare-associated infection</td>
<td>Urinary tract infection that is catheter-associated</td>
<td>SSI</td>
<td></td>
<td>AHRQ, NHSN</td>
</tr>
</tbody>
</table>

### Implementation Guidance

NHSN guidance included on the next few pages.

### Examples
Catheter-Associated Urinary Tract Infection (CAUTI) Event

**Introduction:** The urinary tract is the most common site of healthcare-associated infection, accounting for more than 30% of infections reported by acute care hospitals. Virtually all healthcare-associated urinary tract infections (UTIs) are caused by instrumentation of the urinary tract.

CAUTI can lead to such complications as cystitis, pyelonephritis, gram-negative bacteremia, prostatitis, epididymitis, and orchitis in males and, less commonly, endocarditis, vertebral osteomyelitis, septic arthritis, endophthalmitis, and meningitis in all patients. Complications associated with CAUTI cause discomfort to the patient, prolonged hospital stay, and increased cost and mortality. Each year, more than 13,000 deaths are associated with UTIs.


**Settings:** Surveillance will occur in any inpatient locations where denominator data can be collected, which may include critical intensive care units (ICU), specialty care areas (SCA), stepdown units, and long term care wards. Neonatal units are NOT included. A complete listing of inpatient locations can be found in Chapter 15.

NOTE: It is not required to monitor for CAUTIs after the patient is discharged from the facility, however, if discovered, they should be reported to NHSN. No additional indwelling catheter days are reported.

**Requirements:** Surveillance for CAUTI is performed in at least one inpatient location in the healthcare institution for at least one calendar month as indicated in the *Patient Safety Monthly Reporting Plan* (CDC 57.106).

**Definitions:** As for all infections reported to NHSN, infections associated with complications or extensions of infections already present on admission, unless a change in pathogen or symptoms strongly suggests the acquisition of a new infection area not considered healthcare associated. Therefore, infections that become apparent within the first few days of admission must be carefully reviewed to determine whether they should be considered healthcare associated.

Urinary tract infections (UTI) are defined using symptomatic urinary tract infection (SUTI) criteria or Asymptomatic Bacteremic UTI (ABUTI) criteria (Table 1 and Figure 1). Report UTIs that are catheter-associated (i.e. patient had an indwelling urinary catheter at the time of or within 48 hours before onset of the event).
NOTES:

1. There is no minimum period of time that the catheter must be in place in order for the UTI to be considered catheter-associated. EXAMPLE: Patient has a Foley catheter in place on an inpatient unit. It is discontinued, and 4 days later patient meets the criteria for a UTI. This is not reported as a CAUTI because the time since Foley discontinuation exceeds 48 hours.

2. SUTI 1b and 2b and other UTI (OUTI) cannot be catheter-associated.

Location of attribution: The location where the patient was assigned on the date of the UTI event, which is further defined as the date when the first clinical evidence appeared or the date the specimen use to meet the criterion was collected, whichever came first. EXAMPLE: Patient who had no clinical signs or symptoms of UTI upon arrival to the Emergency Department, has a Foley catheter inserted there before being admitted to the MICU. Within 24 hours of admission to the MICU, patient meets criteria for UTI. This is reported to the NHSN as a CAUTI for the MICU, because the Emergency Department is not an inpatient location and no denominator data are collected there.

TRANSFER RULE EXCEPTION: If a CAUTI develops within 48 hours of transfer from one inpatient location to another in the same facility, or a new facility, the infection is attributed to the transferring location. This is called the Transfer Rule and examples are shown below.

- Patient with a Foley catheter in place in the SICU is transferred to the surgical ward. Thirty six (36) hours later, the patient meets the criteria for UTI. This is reported to NHSN as a CAUTI for the SICU.
- Patient is transferred to the medical ward from the MSICU after having the Foley catheter removed. Within 24 hours, patient meets criteria for a UTI. This is reported to NHSN as a CAUTI for the MSICU.
- Patient with a Foley catheter in place is transferred from the medical ward to the coronary care ICU (CCU). After 4 days in the CCU, the patient meets the criteria for UTI. This is reported to NHSN as a CAUTI for the CCU.
- EXAMPLE: Patient on the urology ward of Hospital A had the Foley catheter removed and is discharged home a few hours later. The ICP from Hospital B calls the next day to report that this patient has been admitted to Hospital B with a UTI. This CAUTI should be reported to NHSN for Hospital A and attributed to the urology ward.

Indwelling catheter: a drainage tube that is inserted into the urinary bladder through the urethra, is left in place, and is connected to a closed collection system; also called a Foley catheter; does not include straight in-and-out catheters.

Numerator Data: The Urinary Tract Infection (UTI) Form (CDC 57.114) is used to collect and report each CAUTI that is identified during the month selected for surveillance. The Instructions for Completion of Urinary Tract Infection Form (Tables of Instructions, Tables 5 and 2a) includes brief instructions for collection and entry of
each data element on the form. The UTI form includes patient demographic information and information on whether or not an indwelling urinary catheter was present. Additional data include the specific criteria met for identifying the UTI, whether the patient developed a secondary bloodstream infection, whether the patient died, and the organisms isolated from cultures and their antimicrobial susceptibilities.

**Denominator data:** Device days and patient days are used for denominators (See Chapter 16 Key Terms). Indwelling urinary catheter days, which are the number of patients with an indwelling urinary catheter device, are collected daily, at the same time each day, according to the chosen location using the appropriate form (CDC 57.117, and 57.118). When denominator data are available from electronic databases, these sources may be used as long as the counts are not substantially different (+/- 5%) from manually collected counts. These daily counts are summed and only the total for the month is entered into NHSN. Indwelling urinary catheter days and patient days are collected separately for each of the locations monitored.

**Data Analyses:** The SIR is calculated by dividing the number of observed infections by the number of expected infections. The number of expected infections, in the context of statistical prediction, is calculated using CAUTI rates from a standard population during a baseline time period as reported in the NHSN Report.

**NOTE:** The SIR will be calculated only if the number of expected HAIs (numExp) is ≥ 1.

\[
\text{SIR} = \frac{\text{Observed (O) HAIs}}{\text{Expected (E) HAIs}}
\]

While the CAUTI SIR can be calculated for single locations, the measure also allows you to summarize your data by multiple locations, adjusting for differences in the incidence of infection among the location types. For example, you will be able to obtain one CAUTI SIR adjusting for all locations reported. Similarly, you can obtain one CAUTI SIR for all specialty care areas in your facility.

The CAUTI rate per 1000 urinary catheter days is calculated by dividing the number of CAUTIs by the number of catheter days and multiplying the result by 1000. The Urinary Catheter Utilization Ratio is calculated by dividing the number of urinary catheter days by the number of patient days. These calculations will be performed separately for the different types of ICUs, specialty care areas, and other locations in the institution, except for neonatal locations.

Table 1: Urinary Tract Infection Criteria

<table>
<thead>
<tr>
<th>Criterion</th>
<th>Urinary Tract Infection (UTI)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Symptomatic Urinary Tract Infection (SUTI)</strong></td>
<td>Must meet at least 1 of the following criteria</td>
</tr>
<tr>
<td>1a</td>
<td>Patient had an indwelling urinary catheter in place at the time of specimen collection and at least 1 of the following signs or symptoms with no other recognized cause: fever (&gt;38°C), suprapubic tenderness, or costovertebral angle pain or tenderness and a positive urine culture of $\geq 10^5$ colony-forming units (CFU)/ml with no more than 2 species of microorganisms.</td>
</tr>
<tr>
<td></td>
<td>---OR-------------------------------------------</td>
</tr>
<tr>
<td></td>
<td>Patient had indwelling urinary catheter removed within the 48 hours prior to specimen collection and at least 1 of the following signs or symptoms with no other recognized cause: fever (&gt;38°C), urgency, frequency, dysuria, suprapubic tenderness, or costovertebral angle pain or tenderness and a positive urine culture of $\geq 10^5$ colony-forming units (CFU)/ml with no more than 2 species of microorganisms.</td>
</tr>
<tr>
<td>1b</td>
<td>Patient did not have an indwelling urinary catheter in place at the time of specimen collection nor within 48 hours prior to specimen collection and has at least 1 of the following signs or symptoms with no other recognized cause: fever (&gt;38°C) in a patient that is ≤65 years of age, urgency, frequency, dysuria, suprapubic tenderness, or costovertebral angle pain or tenderness and a positive urine culture of $\geq 10^5$ CFU/ml with no more than 2 species of microorganisms.</td>
</tr>
<tr>
<td>2a</td>
<td>Patient had an indwelling urinary catheter in place at the time of specimen collection and at least 1 of the following signs or symptoms with no other recognized cause: fever (&gt;38°C), suprapubic tenderness, or costovertebral angle pain or tenderness and a positive urinalysis demonstrated by at least 1 of the following findings: a. positive dipstick for leukocyte esterase and/or nitrite b. pyuria (urine specimen with $\geq 10$ white blood cells [WBC]/mm$^3$ of unspun urine or $\geq 3$ WBC/high power field of spun urine)</td>
</tr>
<tr>
<td>Criterion</td>
<td>Urinary Tract Infection (UTI)</td>
</tr>
<tr>
<td>-----------</td>
<td>------------------------------</td>
</tr>
</tbody>
</table>
| c. microorganisms seen on Gram stain of unspun urine
and
a positive urine culture of $\geq 10^3$ and $<10^5$ CFU/ml with no more than 2 species of microorganisms. | |

Patient had indwelling urinary catheter **removed within the 48 hours prior** to specimen collection
and
at least 1 of the following signs or symptoms with no other recognized cause:
fever (>38°C), urgency, frequency, dysuria, suprapubic tenderness, or costovertebral angle pain or tenderness
and
a positive urinalysis demonstrated by at least 1 of the following findings:
   a. positive dipstick for leukocyte esterase and/or nitrite
   b. pyuria (urine specimen with $\geq 10$ white blood cells [WBC]/mm$^3$ of unspun urine or $\geq 3$ WBC/high power field of spun urine)
   c. microorganisms seen on Gram stain of unspun urine
and
a positive urine culture of $\geq 10^3$ and $<10^5$ CFU/ml with no more than 2 species of microorganisms.

2b Patient did **not** have an indwelling urinary catheter in place at the time of specimen collection nor within 48 hours prior to specimen collection
and
has at least 1 of the following signs or symptoms with no other recognized cause:
fever (>38°C) in a patient that is $\leq 65$ years of age, urgency, frequency, dysuria, suprapubic tenderness, or costovertebral angle pain or tenderness
and
a positive urinalysis demonstrated by at least 1 of the following findings:
   a. positive dipstick for leukocyte esterase and/or nitrite
   b. pyuria (urine specimen with $\geq 10$ WBC/mm$^3$ of unspun urine or $\geq 3$ WBC/high power field of spun urine)
   c. microorganisms seen on Gram stain of unspun urine
and
a positive urine culture of $\geq 10^3$ and $<10^5$ CFU/ml with no more than 2 species of microorganisms.

3 Patient $\leq 1$ year of age with or without an indwelling urinary catheter has at least 1 of the following signs or symptoms with no other recognized cause: fever (>38°C core), hypothermia (<36°C core), apnea, bradycardia, dysuria, lethargy, or vomiting
### Urinary Tract Infection (UTI)

<table>
<thead>
<tr>
<th>Criterion</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>4</td>
<td>Patient ≤1 year of age with or without an indwelling urinary catheter has at least 1 of the following signs or symptoms with no other recognized cause: fever (&gt;38°C core), hypothermia (&lt;36°C core), apnea, bradycardia, dysuria, lethargy, or vomiting and a positive urinalysis demonstrated by at least one of the following findings: a. positive dipstick for leukocyte esterase and/or nitrite b. pyuria (urine specimen with ≥10 WBC/mm³ of unspun urine or ≥3 WBC/high power field of spun urine) c. microorganisms seen on Gram’s stain of unspun urine and a positive urine culture of between ≥10³ and &lt;10⁵ CFU/ml with no more than two species of microorganisms.</td>
</tr>
</tbody>
</table>

### Asymptomatic Bacteremic Urinary Tract Infection (ABUTI)

<table>
<thead>
<tr>
<th>Criterion</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>4</td>
<td>Patient with or without an indwelling urinary catheter has no signs or symptoms (i.e., for any age patient, no fever (&gt;38°C), urgency, frequency, dysuria, suprapubic tenderness, or costovertebral angle pain or tenderness, OR for a patient ≤1 year of age, no fever (&gt;38°C core), hypothermia (&lt;36°C core), apnea, bradycardia, dysuria, lethargy, or vomiting) and a positive urine culture of &gt;10⁵ CFU/ml with no more than 2 species of uropathogen microorganisms* and a positive blood culture with at least 1 matching uropathogen microorganism to the urine culture, or at least 2 matching blood cultures drawn on separate occasions if the matching pathogen is a common skin contaminant.</td>
</tr>
</tbody>
</table>

---

**Comments**

- Urinary catheter tips should not be cultured and are not acceptable for the diagnosis of a urinary tract infection.
- Urine cultures must be obtained using appropriate technique, such as clean catch collection or catheterization. Specimens from indwelling catheters should be aspirated through the disinfected sampling ports.
- In infants, urine cultures should be obtained by bladder catheterization or suprapubic aspiration; positive urine cultures from bag specimens are unreliable and should be confirmed by specimens aseptically obtained by catheterization or suprapubic aspiration.

* Uropathogen microorganisms are: Gram-negative bacilli, *Staphylococcus* spp., yeasts, beta-hemolytic *Streptococcus* spp., *Enterococcus* spp., *G. vaginalis*, *Aerococcus urinae*, and *Corynebacterium* (urease positive).
### Device-associated Module

**CAUTI**

<table>
<thead>
<tr>
<th>Criterion</th>
<th>Urinary Tract Infection (UTI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>- Urine specimens for culture should be processed as soon as possible, preferably within 1 to 2 hours. If urine specimens cannot be processed within 30 minutes of collection, they should be refrigerated, or inoculated into primary isolation medium before transport, or transported in an appropriate urine preservative. Refrigerated specimens should be cultured within 24 hours.</td>
<td></td>
</tr>
<tr>
<td>- Urine specimen labels should indicate whether or not the patient is symptomatic.</td>
<td></td>
</tr>
<tr>
<td>- Report secondary bloodstream infection = “Yes” for all cases of Asymptomatic Bacteremic Urinary Tract Infection (ABUTI).</td>
<td></td>
</tr>
<tr>
<td>- Report only pathogens in both blood and urine specimens for ABUTI.</td>
<td></td>
</tr>
<tr>
<td>- Report <em>Corynebacterium</em> (urease positive) as either <em>Corynebacterium</em> species unspecified (COS) or, as <em>C. urealyticum</em> (CORUR) if so speciated.</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Criterion</th>
<th>Other Urinary Tract Infection (OUTI) (kidney, ureter, bladder, urethra, or tissue surrounding the retroperineal or perinephric space)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Other infections of the urinary tract must meet at least 1 of the following criteria:</td>
<td></td>
</tr>
<tr>
<td>1 Patient has microorganisms isolated from culture of fluid (other than urine) or tissue from affected site.</td>
<td></td>
</tr>
<tr>
<td>2 Patient has an abscess or other evidence of infection seen on direct examination, during a surgical operation, or during a histopathologic examination.</td>
<td></td>
</tr>
<tr>
<td>3 Patient has at least 2 of the following signs or symptoms with no other recognized cause: fever (&gt;38°C), localized pain, or localized tenderness at the involved site and at least 1 of the following:</td>
<td></td>
</tr>
<tr>
<td>a. purulent drainage from affected site</td>
<td></td>
</tr>
<tr>
<td>b. microorganisms cultured from blood that are compatible with suspected site of infection</td>
<td></td>
</tr>
<tr>
<td>c. radiographic evidence of infection (e.g., abnormal ultrasound, CT scan, magnetic resonance imaging [MRI], or radiolabel scan [gallium, technetium]).</td>
<td></td>
</tr>
<tr>
<td>4 Patient &lt; 1 year of age has at least 1 of the following signs or symptoms with no other recognized cause: fever (&gt;38°C core), hypothermia (&lt;36°C core), apnea, bradycardia, lethargy, or vomiting and at least 1 of the following:</td>
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<tr>
<td>a. purulent drainage from affected site</td>
<td></td>
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<td>b. microorganisms cultured from blood that are compatible with suspected site of infection</td>
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<tr>
<td>c. radiographic evidence of infection, (e.g., abnormal ultrasound, CT scan, magnetic resonance imaging [MRI], or radiolabel scan [gallium, technetium]).</td>
<td></td>
</tr>
</tbody>
</table>

| Comment | - Report infections following circumcision in newborns as SST-CIRC. |
Figure 1: Identification and Categorization of SUTI Indwelling Catheter at the Time of Specimen Collection

Patient had an indwelling urinary catheter at the time of specimen collection

At least 1 of the following with no other recognized cause:
- fever (>38°C)
- suprapubic tenderness
- costovertebral angle pain or tenderness

Signs and Symptoms

Urinalysis

A positive urinalysis demonstrated by at least 1 of the following findings:
- positive dipstick for leukocyte esterase and/or nitrite
- pyuria (urine specimen with ≥10 WBC/mm³ of unspun urine or ≥3 WBC/high power field of spun urine)
- microorganisms seen on Gram stain of unspun urine

Culture Evidence

A positive urine culture of ≥10⁵ CFU/ml with no more than 2 species of microorganisms

SUTI – Criterion 1a

CAUTI

A positive urine culture of ≥10³ and <10⁵ CFU/ml with no more than 2 species of microorganisms

SUTI – Criterion 2a

CAUTI
Figure 2: Identification and Categorization of SUTI Indwelling Catheter Discontinued in Prior 48 Hours

Patient had an indwelling urinary catheter discontinued within 48 hours prior to specimen collection

At least 1 of the following with no other recognized cause:
- fever (>38°C)
- urgency
- frequency
- dysuria
- suprapubic tenderness
- costovertebral angle pain or tenderness

A positive urinalysis demonstrated by at least 1 of the following findings:
- positive dipstick for leukocyte esterase and/or nitrite
- pyuria (urine specimen with ≥10 WBC/mm² of unspun urine or ≥3 WBC/high power field of spun urine)
- microorganisms seen on Gram stain of unspun urine

A positive urine culture of ≥10⁵ CFU/ml with no more than 2 species of microorganisms

SUTI—Criterion 1a

CAUTI

A positive urine culture of ≥10³ and <10⁵ CFU/ml with no more than 2 species of microorganisms

SUTI—Criterion 2a

CAUTI
Figure 3: Identification and Categorization of SUTI Without Indwelling Catheter at Time of or Within 48 Hours Prior to Specimen Collection

Patient did not have an indwelling urinary catheter at the time of specimen collection nor within 48 hours prior to specimen collection.

**Signs and Symptoms**
- At least 1 of the following with no other recognized cause:
  - Fever (>38°C) in a patient that is ≤65 years of age
  - Urinary urgency
  - Frequent urination
  - Dysuria
  - Suprapubic tenderness
  - Costovertebral angle pain or tenderness

**Urinalysis**
- A positive urinalysis demonstrated by at least 1 of the following findings:
  - Positive dipstick for leukocyte esterase and/or nitrite
  - Pyuria (urine specimen with ≥10 WBC/mm³ of unspun urine or ≥3 WBC/high power field of spun urine)
  - Microorganisms seen on Gram stain of unspun urine

**Culture Evidence**
- A positive urine culture of ≥10^6 CFU/ml with no more than 2 species of microorganisms
- A positive urine culture of ≥10^3 and < 10^6 CFU/ml with no more than 2 species of microorganisms

SUTI – Criterion 1b
SUTI – Criterion 2b
Figure 4: Identification and Categorization of SUTI in Patient ≤1 Year of Age

Patient ≤1 year of age (with or without an indwelling urinary catheter)

At least 1 of the following with no other recognized cause:
- fever (>38°C core)
- dysuria
- hypothermia (<36°C core)
- lethargy
- apnea
- vomiting
- bradycardia

Urinalysis

A positive urinalysis demonstrated by at least 1 of the following findings:
- positive dipstick for leukocyte esterase and/or nitrite
- pyuria (urine specimen with ≥10 WBC/mm³ of unspun urine or ≥3 WBC/high power field of spun urine)
- microorganisms seen on Gram stain of unspun urine

Culture Evidence

A positive urine culture of ≥10⁶ CFU/ml with no more than 2 species of microorganisms

SUTI — Criterion 3

Was an indwelling urinary catheter in place within the last 48 hours?

- Yes
- No

- CAUTI
- SUTI

A positive urine culture of <10³ and <10⁶ CFU/ml with no more than 2 species of microorganisms

SUTI — Criterion 4

Was an indwelling urinary catheter in place within the last 48 hours?

- Yes
- No

- CAUTI
- SUTI
Figure 5: Identification of Asymptomatic Bacteremic Urinary Tract Infection (ABUTI)

Patient with or without an indwelling urinary catheter

**Patient of any age**
- NONE of the following:
  - fever (>38°C)
  - urgency
  - frequency
  - dysuria
  - suprapubic pain
  - costovertebral angle pain or tenderness

**Patient ≤1 year of age**
- NONE of the following:
  - fever (>38°C core)
  - hypothermia (<36°C core)
  - apnea
  - bradycardia
  - lethargy
  - vomiting

A positive urine culture of ≥10⁶ CFU/ml with no more than 2 species of microorganisms *

A positive blood culture with at least 1 matching uropathogen microorganism* to the urine culture

Asymptomatic Bacteremic Urinary Tract Infection (ABUTI)

*Uropathogen microorganisms are: Gram-negative bacilli, *Staphylococcus* spp., yeasts, beta-hemolytic *Streptococcus* spp., *Enterococcus* spp., *G. vaginalis*, *Aerococcus urinae*, *Corynebacterium* (urease positive)*.

†Report *Corynebacterium* (urease positive) as either *Corynebacterium species unspecified* (COS) or, as *C. urealyticum* (CORUR) if so speciated.
<table>
<thead>
<tr>
<th>event category</th>
<th>full term</th>
<th>short term</th>
<th>specifications</th>
<th>standard</th>
</tr>
</thead>
<tbody>
<tr>
<td>other</td>
<td>healthcare-associated infection</td>
<td>HAI</td>
<td></td>
<td>NRS</td>
</tr>
</tbody>
</table>

**Implementation guidance**

This area is intended to capture events not previously covered in above categories. Reviewing 2009-2010 data categorical inclusion: *E Coli* in sputum on ventilator, stool infection, necrotizing pancreatitis.

**Examples**

*Actual sentinel event:* A 68 year-old male is admitted with stool impaction and a urinary tract infection. After 4 days in hospital, the patient’s condition worsens, and he is subsequently found to have *C difficile* in his stool. The infection results in sepsis and 3-day prolonged hospitalization with admission to intensive care unit.
OTHER

other – specify

<table>
<thead>
<tr>
<th>event category</th>
<th>full term</th>
<th>short term</th>
<th>specifications</th>
<th>standard</th>
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<td>other</td>
<td>specifications</td>
<td>NRS</td>
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</tbody>
</table>

implementation guidance

Any unexpected death not elsewhere classified qualifies as a sentinel event and should be reported under the other category with a brief description accompanying it.

examples

actual sentinel event: A patient undergoes a surgical procedure and pre-operative assessment reveals no allergies. The patient receives intravenous Cephalexin and has a subsequent anaphylactic reaction. All attempts are made to revive patient, but they are unsuccessful, and the patient dies.

actual sentinel event: A 62 year-old patient with underlying HTN, diabetes, and obesity is medically cleared for a hip replacement. The patient is anesthetized without complication. An intra-operative cardiac arrhythmia is noted, and the patient is defibrillated without success and is subsequently pronounced dead.

not a sentinel event: 72 year-old Hispanic male diagnosed with end-stage pancreatic cancer agrees to a palliative surgery to improve comfort. The day after surgery, the patient has a cardiac arrhythmia and dies. Death is attributed to the patient’s terminal cancer. This does not need to be reported.
<table>
<thead>
<tr>
<th>term</th>
<th>definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>abduction</td>
<td>means the taking away of a person by persuasion, by fraud, or by open force or violence. It includes convincing someone, particularly a minor or a woman he/she is better off leaving with the persuader, telling the person he/she is needed, or that the mother or father wants him/her to come with the abductor.</td>
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<tr>
<td>adverse</td>
<td>describes a consequence of care that results in an undesired outcome. It does not address preventability.</td>
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<tr>
<td>associated with</td>
<td>means that it is reasonable to initially assume that the adverse event was due to the referenced course of care; further investigation and/or root cause analysis of the unplanned event may be needed to confirm or refute the presumed relationship.</td>
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<tr>
<td>authorized</td>
<td>means the guardian or other individual(s) having the legally recognized ability to consent on behalf of a minor or incapacitated individual (surrogate), or person designated by the surrogate to release or consent for the patient.</td>
</tr>
<tr>
<td>decision-making capacity</td>
<td>is the ability to understand information relevant to a decision and the ability to appreciate the reasonably foreseeable consequences of a decision (or lack of a decision).</td>
</tr>
<tr>
<td>deep tissue injury</td>
<td>presents as a purple or maroon localized area of discolored intact skin or blood-filled blister due to damage of underlying soft tissue from pressure and/or shear. The area may be preceded by tissue that is painful, firm, mushy, boggy, warmer or cooler as compared to adjacent tissue.</td>
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<tr>
<td>device</td>
<td>See Medical Device.</td>
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<tr>
<td>elopement</td>
<td>refers to a situation where a patient or resident who is cognitively, physically, mentally, emotionally, and/or chemically impaired wanders/walks/runs away, escapes, or otherwise leaves a caregiving institution or setting unsupervised, unnoticed, and/or prior to their scheduled discharge.</td>
</tr>
<tr>
<td>event</td>
<td>means a discrete, auditable, and clearly defined occurrence.</td>
</tr>
<tr>
<td>healthcare setting</td>
<td>means any facility or office, including a discrete unit of care within such facility, that is organized, maintained, and operated for the diagnosis, prevention, treatment, rehabilitation, convalescence or other care of human illness or injury, physical or mental, including care during and after pregnancy. Healthcare settings include, but are not limited to, hospitals, nursing homes, rehabilitation centers, medical centers, office-based practices, outpatient dialysis centers, reproductive health centers, independent clinical laboratories, hospices, ambulatory surgical centers, and pharmacies. The boundary of a healthcare setting (the “grounds”) is the physical area immediately adjacent to the setting’s main buildings. It does not include nonmedical</td>
</tr>
<tr>
<td><strong>high alert medications</strong></td>
<td>are those medications that have a high risk of causing serious injury or death to a patient if they are misused. Examples of high-alert medications include anticoagulants and IV antithrombotics, insulin, cytotoxic chemotherapy, concentrated electrolytes, IV digoxin, opiate narcotics, neuromuscular blocking agents, and adrenergic agonists. <em>The recommended “High Alert Medication List” is available at the Institute for Safe Medication Practices’ website, <a href="http://www.ismp.org">www.ismp.org</a>.</em></td>
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<tr>
<td><strong>infant</strong></td>
<td>is a child under the age of one year. (SRE 2006; Stedman’s online dictionary)</td>
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<tr>
<td><strong>informed consent</strong></td>
<td>involves a process of shared decisionmaking in which discussion between a person who would receive a treatment, including surgery or invasive procedure, and the caregiver/professional person who explains the treatment, provides information about possible benefits, risks and alternatives, and answers questions that result in the person’s authorization or agreement to undergo a specific medical intervention. Documentation of this discussion should result in an accurate and meaningful entry in the patient record, which could include a signed “consent form.” Signing a consent form does not constitute informed consent; it provides a record of the discussion.</td>
</tr>
<tr>
<td><strong>injury</strong></td>
<td>as used in this report has a broad meaning. It includes physical or mental damage that substantially limits one or more of the major life activities of an individual in the short term, which may become a disability if extended long term. Further, injury includes a substantial change in the patient’s long-term risk status such that care or Appendix B - Glossary B-3 National Quality Forum monitoring, based on accepted national standards, is required that was not required before the event. <em>(Of note, states and other entities may use alternate definitions for the term “disability.”)</em></td>
</tr>
<tr>
<td><strong>largely preventable</strong></td>
<td>recognizes that some of the events on the SRE list are not universally avoidable, given the complexity of healthcare and current knowledge.</td>
</tr>
<tr>
<td><strong>low-risk pregnancy</strong></td>
<td>refers to a woman aged 18-39, with no previous diagnosis of essential hypertension, renal disease, collagen-vascular disease, liver disease, cardiovascular disease, placenta previa, multiple gestation, intrauterine growth retardation, smoking, pregnancy-induced hypertension, premature rupture of membranes, or other previously documented condition that poses a high risk of poor pregnancy outcome.</td>
</tr>
<tr>
<td><strong>medical device</strong></td>
<td>is an instrument, apparatus, implement, machine, contrivance, implant, in vitro reagent, or other similar or related article, including a component part, or accessory, which is recognized in the official National Formulary, or the United States Pharmacopoeia, or any supplement to them; intended for use in the diagnosis of disease or other conditions, or in the cure, mitigation, treatment, or prevention of disease, in man or other animals; or intended to affect the structure or any function of the body of man or other animals, and which does not achieve any of its primary intended purposes through chemical action within or on the body of man or other animals and which is not dependent upon being metabolized for the achievement of any of its primary intended purposes.¹</td>
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<tr>
<td><strong>medication error</strong></td>
<td>means any preventable event that may cause or lead to inappropriate medication use or patient harm while the medication is in the control of the healthcare professional, patient, or consumer. Such events may be related to professional practice, healthcare products, procedures, and systems, including prescribing; order communication; product labeling, packaging and nomenclature; compounding; dispensing; distribution; administration; education; monitoring; and use.²</td>
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<tr>
<td><strong>neonate</strong></td>
<td>is a newborn less than 28 days of age.</td>
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<td><strong>patient</strong></td>
<td>means a person who is a recipient of healthcare. A person becomes a patient at the point that they are being “cared for” in the facility. Being “cared for” begins when they are first engaged by a member of the care team, e.g. assessment by the triage nurse in the E.D., walking with the phlebotomist to the lab for a lab draw. A patient is no longer considered a patient at the point that they are no longer under the care of a member of the care team, e.g. the nursing assistant has safely assisted the patient to the car from an inpatient stay; the ambulating patient that does not need assistance leaves the radiology department following an outpatient test.³</td>
</tr>
<tr>
<td><strong>pressure ulcer, stage 3</strong></td>
<td>is defined as full thickness tissue loss. Subcutaneous fat may be visible, but bone, tendon, or muscle is not exposed. Slough may be present. May include undermining and tunneling. The depth of a Stage 3 pressure ulcer varies by anatomical location. The bridge of the nose, ear, occiput, and malleolus do not have subcutaneous tissue and Stage 3 ulcers can be shallow. In contrast, areas of significant adiposity can develop extremely deep Stage 3 pressure ulcers. Bone/tendon is not visible or directly palpable.⁴</td>
</tr>
<tr>
<td><strong>pressure ulcer, stage 4</strong></td>
<td>is defined as full thickness tissue loss with exposed bone, tendon, or muscle. Slough or eschar may be present. Often includes undermining and tunneling. The depth of a Stage 4 pressure ulcer varies by anatomical location. The bridge of the nose, ear, occiput, and malleolus do not have subcutaneous tissue and these ulcers can be shallow. Stage 4 ulcers can extend into muscle and/or supporting structures (e.g., fascia, tendon, or joint capsule) making osteomyelitis or osteitis likely to occur. Exposed</td>
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<td><strong>Gbential Event Reporting Guidance</strong></td>
<td><strong>Glossary</strong></td>
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<tr>
<td><strong>bone/tendon is visible or directly palpable.</strong>&lt;sup&gt;4&lt;/sup&gt;</td>
<td><strong>pressure ulcer, unstageable</strong> is defined as full thickness tissue loss in which the actual depth of the ulcer is completely obscured by slough and/or eschar in the wound bed. Until enough slough and/or exchar are removed to expose the base of the wound, the true depth cannot be determined; but it will be either Stage 3 or Stage 4.&lt;sup&gt;4&lt;/sup&gt;</td>
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<tr>
<td><strong>preventable</strong> describes an event that could have been anticipated and prepared for, but that occurs because of an error or other system failure.</td>
<td><strong>restraints</strong> is defined by The Joint Commission, the Centers for Medicare &amp; Medicaid Services, and by some states. The appropriate source(s) should be consulted for the definition required by the setting and/or jurisdiction in which a presumptive event occurs. In the event none of those definitions apply to an institution, the following definition, which is intended to capture definitions from the named organizations, is offered: Restraints means any method of restricting a patient’s freedom of movement that is not a usual and customary part of a medical diagnostic or treatment procedure to which the patient or his or her legal representative has consented; is not indicated to treat the patient’s medical condition or symptoms; or does not promote the patient’s independent functioning.</td>
</tr>
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<td><strong>serious</strong> describes an event that can result in death, loss of a body part, disability, loss of bodily function, or require major intervention for correction (e.g., higher level of care, surgery).</td>
<td><strong>sexual abuse</strong> NRS 200.366 Sexual assault: Definition; penalties. 1. A person who subjects another person to sexual penetration, or who forces another person to make a sexual penetration on himself or herself or another, or on a beast, against the will of the victim or under conditions in which the perpetrator knows or should know that the victim is mentally or physically incapable of resisting or understanding the nature of his or her conduct, is guilty of sexual assault.</td>
</tr>
<tr>
<td><strong>surgery</strong> NAC 449.9743 “Surgery” defined. (NRS 449.037) “Surgery” means the treatment of a human being by a physician using one or more of the following procedures: 1. Cutting into any part of the body using a scalpel, electrocautery or any other means for diagnosis or the removal or repair of diseased or damaged tissue, organs, tumors or foreign bodies. 2. The reduction of a fracture or the dislocation of a bone, joint or bony structure. 3. The repair of a malformation of the body resulting from an injury, a birth defect or another cause, that requires cutting and manipulation or a suture. 4. An instrumentation of the uterine cavity of a woman for diagnostic or therapeutic purposes, including the procedure commonly known as dilation and curettage. 5. Any instrumentation of, or injection of a substance into, the uterine cavity of a woman to terminate a pregnancy. 6. Any procedure to sterilize a human being.</td>
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<td><strong>unambiguous</strong></td>
<td>refers to an event that is clearly defined and easily identified.</td>
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<td><strong>unintended retention</strong></td>
<td>of a foreign object refers to a foreign object introduced into the body during a surgical or other invasive procedure, without removal prior to the end of the surgery or procedure, which the surgeon or other practitioner did not intend to leave in the body.</td>
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</tbody>
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