National Healthcare Safety Network (NHSN)
Outpatient Procedure Component Manual

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Please Note: The NHSN Outpatient Procedure Component Manual is updated annually based on subject matter expert review and user feedback. Over time, certain chapters may be retired or moved to another NHSN component. To avoid confusion, the chapters in the OPC manual do not shift to account for these changes.
National Healthcare Safety Network (NHSN) Overview

The NHSN is a secure, Internet-based surveillance system that expands and integrates patient and healthcare personnel safety surveillance systems managed by the Division of Healthcare Quality Promotion (DHQP) at the Centers for Disease Control and Prevention. In addition, facilities that participate in certain reporting programs operated by the Centers for Medicare and Medicaid Services (CMS) can do so through use of NHSN. Furthermore, some U.S. states use NHSN as a means for healthcare facilities to submit data on healthcare-associated infections (HAIs) and transfusion-related adverse events mandated through their specific state legislation.

NHSN enables healthcare facilities to collect and use data about HAIs, adherence to clinical practices known to prevent HAIs, the incidence or prevalence of multidrug-resistant organisms within their organizations, trends and coverage of healthcare personnel safety and vaccination, and adverse events related to the transfusion of blood and blood products.

The NHSN includes six components: Patient Safety, Long-term Care Facility, Outpatient Dialysis, Healthcare Personnel Safety, Biovigilance, and Outpatient Procedure (Figure 1).

**Figure 1: NHSN Components**
The **Patient Safety Component** includes five modules that focus on events associated with medical devices, surgical procedures, antimicrobial agents used during healthcare, multidrug resistant organisms, and Coronavirus Infectious Disease 2019 (COVID-19).

- **Device-associated Module:**
  - Bloodstream Infection (CLABSI – Central line-associated bloodstream infection)
  - Central line insertion practices (CLIP) adherence
  - Urinary Tract Infection (CAUTI – Catheter-associated urinary tract infection)
  - Pediatric Ventilator-associated events (PedVAE) (NICU and pediatric locations only)
  - Ventilator-associated events (VAE) (adult locations only)
  - Pneumonia (VAP – Ventilator-associated pneumonia) - in pediatric locations (in-plan* or off-plan*), or NICU and adult locations (off-plan* only)

- **Procedure-associated Module:**
  - Surgical Site Infection (SSI)

- **Antimicrobial Use and Resistance Module (AUR)**

- **Multidrug-Resistant Organism and *Clostridium difficile* Infection (MDRO/CDI) Module**

- **Coronavirus Infectious Disease 2019 (COVID-19) Module (off plan* only):**
  - Patient Impact and Hospital Capacity Pathway
  - Healthcare Worker Staffing Pathway
  - Healthcare Supply Pathway

*Note:* “In-plan” surveillance means that the facility has committed to following the NHSN surveillance protocol, in its entirety, for that particular event, as shown in the facility’s NHSN monthly reporting plan. “Off-plan” surveillance is surveillance that is done because a facility has decided to track a particular event for internal use. Data that are entered into NHSN “off-plan” are not included in NSHN annual reports or other NHSN publications. A facility makes no commitment to follow the NHSN protocol for “off-plan” events. Further, “off-plan” data cannot be uploaded into NHSN via Clinical Document Architecture (CDA) and must be manually entered. Instructions and standardized surveillance methods and definitions for each module of the Patient Safety Component are provided in this manual and on the NHSN website ([www.cdc.gov/nhsn](http://www.cdc.gov/nhsn)). Modules may be used singly or simultaneously.

The NHSN **Long-term Care Facility Component** provides long-term care facilities (LTCFs) with standardized surveillance methods and definitions for four modules: (1) Multidrug resistant organism (MDRO) and *Clostridioides difficile* Infection (CDI) laboratory-identified (LabID) Events; (2) Urinary Tract Infections (UTI); (3) Prevention Process Measures; and (4) Coronavirus Infectious Disease (COVID-19). The component is ideal for use by nursing homes, skilled nursing facilities, chronic care facilities, and assisted living and residential care facilities. LTCF surveillance protocols, training materials, data collection forms, instructions, and other supporting materials are provided on the Long-term Care Facility Component website: [https://www.cdc.gov/nhsn/ltc/index.html](https://www.cdc.gov/nhsn/ltc/index.html).

Outpatient hemodialysis centers have several surveillance options tailored to their patients and setting in the **Dialysis Component**. The component consists of 3 modules: 1) Dialysis Event; 2) Prevention Process Measures; and (3) Dialysis Patient Influenza Vaccination. Facilities that treat hemodialysis outpatients should refer to the Dialysis Component instructions and standardized surveillance methods and definitions at [www.cdc.gov/nhsn/dialysis/index.html](http://www.cdc.gov/nhsn/dialysis/index.html).
There are two modules in the **Healthcare Personnel Safety (HPS) Component** of NHSN: The Healthcare Personnel Exposure Module and the Healthcare Personnel Vaccination Module. The Healthcare Personnel Exposure Module includes: Blood/Body Fluid Exposure Only; Blood/Body Fluid Exposure with Exposure Management; and Influenza Exposure Management. This module is no longer available for enrollment and should only be used by facilities that have already been reporting Blood/Body Fluid Exposure and Exposure Management data to the system. The Healthcare Personnel Vaccination Module includes: Influenza Vaccination Summary. Data collected in this surveillance system can assist healthcare facilities, health systems, and public health agencies to monitor and report trends in blood/body fluid exposures, to characterize antiviral medication use for exposures to influenza, and to monitor influenza vaccination coverage among healthcare personnel. These modules may be used separately or simultaneously. Instructions and standardized surveillance methods and definitions for the Healthcare Personnel Vaccination Module is provided in the NHSN Manual: HPS Component Protocol [https://www.cdc.gov/nhsn/pdfs/hps-manual/vaccination/hps-flu-vaccine-protocol.pdf](https://www.cdc.gov/nhsn/pdfs/hps-manual/vaccination/hps-flu-vaccine-protocol.pdf)

The **NHSN Biovigilance Component**, Hemovigilance Module facilitates national surveillance of transfusion-related recipient adverse events. The Hemovigilance Module is designed for transfusion service staff to collect data on annual facility and transfusion service characteristics, individual reports on adverse transfusion reactions, errors or accidents associated with adverse reactions, and monthly counts of transfused or discarded components. The Hemovigilance Module surveillance protocol, training materials, data collection forms, instructions, and other supporting materials are provided on the Hemovigilance Module website: [www.cdc.gov/nhsn/acute-care-hospital/bio-hemo/index.html](http://www.cdc.gov/nhsn/acute-care-hospital/bio-hemo/index.html).

The **Outpatient Procedure Component (OPC)** includes two modules that focus on adverse events associated with surgical procedures performed in Ambulatory Surgery Centers (ASCs). The two modules include Same Day Outcome Measures and Surgical Site Infections.

- **Same Day Outcome Measures (OPC-SDOM)** are a grouping of outpatient care quality indicators that represent a broad range of risks encountered by patients accessing care in various outpatient settings. The four individual outcome measures are:
  - Patient Burn
  - Patient Fall
  - Wrong Site, Wrong Side, Wrong Patient, Wrong Procedure, Wrong Implant
  - All-Cause Hospital Transfer/Admission

- **Surgical Site Infection (OPC-SSI)** - SSI surveillance for outpatient operative procedures using the Outpatient Procedure Component (OPC) replaces the use of the Patient Safety Component SSI event chapter for ASCs.

The OPC surveillance protocols, training materials, data collection forms, instructions, and other supporting materials are provided on the Outpatient Procedure Component website: [https://www.cdc.gov/nhsn/ambulatory-surgery/index.html](https://www.cdc.gov/nhsn/ambulatory-surgery/index.html).
Surveillance Techniques

Some of the options in the following modules require active, patient-based, prospective surveillance of events and their corresponding denominator data by a trained Infection Preventionist (IP). This means that the IP shall seek out infections during a patient’s stay by screening a variety of data sources, such as laboratory, pharmacy, admission/discharge/transfer, radiology/imaging, and pathology databases, as well as patient charts, including history and physical exam notes, nurses’/physicians’ notes, temperature charts, etc. Others may be trained to screen data sources for these infections, but the IP must make the final determination. Laboratory-based surveillance should not be used alone, unless all possible criteria for identifying an infection are solely determined by laboratory evidence (for example, LabID event detection in the MDRO/CDI Module). Retrospective chart reviews should be used only when patients are discharged before all information can be gathered. NHSN forms should be used to collect all required data, using the NHSN definitions of each data field. To minimize the IP’s data collection burden, others may be trained to collect the denominator data and process of care data (for example, central line insertion practices).

Procedure-Associated Module

Surgical site infection (SSI) monitoring is offered through this module. SSI surveillance requires active, patient-based, prospective surveillance techniques (see Surveillance Techniques above). To minimize IPs’ workload of collecting denominator data, operating room data may be downloaded (see file specifications at: https://www.cdc.gov/nhsn/pdfs/ps-analysis-resources/ImportingProcedureData.pdf)

Both pre-discharge and post-discharge surveillance methods should be used to detect SSIs. Surveillance may include both inpatient and outpatient operative procedures. These methods include 1) direct examination of patients’ wounds during hospitalization, or follow-up visits to either surgery clinics or physicians’ offices, 2) review of medical records or surgery clinic patient records, 3) surgeon surveys by mail or telephone, and 4) patient surveys by mail or telephone (though patients may have a difficult time assessing their infections). Any combination of these methods is acceptable for use; however, CDC criteria for SSI must be applied.

Device-Associated Module

Medical instrumentation increases the risk of development of an HAI and most patients admitted for health care are exposed to some kind of medical device in the course of their treatment. Such devices include, but are not limited to, vascular and urinary catheters, and ventilators. NHSN enables facilities to monitor infectious complications associated with the use of these devices and to monitor processes related to their use which might increase infection risk. Specifically, surveillance of central line-associated bloodstream infection (CLABSI), catheter-associated urinary tract infection (CAUTI), ventilator-associated events (VAE), and/or ventilator-associated pneumonia (VAP) is possible using the NHSN. In addition, central line insertion practices (CLIP) can be monitored to inform facilities of the appropriateness of their processes and how they may relate to HAI development. See Dialysis Component for detailed instructions for Dialysis Event (DE) surveillance of hemodialysis outpatients (www.cdc.gov/nhsn/dialysis/index.html).
Device-associated denominator data should be collected at the same time each day, or by weekly sampling methods, in certain locations, for CLABSI and CAUTI surveillance (see the CLABSI and CAUTI protocols for guidance). When denominator data are available from electronic databases (for example, ventilator days from respiratory therapy), these sources may be used as long as the counts are not substantially different (+/- 5%) from manually-collected counts that have been validated for a minimum of three months. See the respective device-associated event protocols for detailed surveillance instructions.

**Antimicrobial Use and Resistance (AUR) Module**

The use of antimicrobial agents has a direct effect on antimicrobial resistance patterns of pathogens. The observed increase in multidrug resistance is in part due to inappropriate prescription of, as well as only partial completion of courses of antibiotics.

The AUR Module allows facilities to collect information on the amount of antimicrobials that are used for patient care within their systems, as well as to collect data on the prevalence of drug-resistant organisms in their inpatient and outpatient areas. Electronic capture and reporting of microbiology and pharmacy data are the only available options for reporting data into this module.

See the [Antimicrobial Use and Resistance](#) protocol for detailed surveillance instructions.

**Multidrug-resistant Organism and Clostridium difficile Infection (MDRO/CDI) Module**

The NHSN MDRO/CDI Module offers a means for facilities to meet criteria and metrics that are outlined in several organizational guidelines to control and measure the spread of MDROs and CDI within their healthcare system. The module has two separate and independent reporting options, Laboratory-identified (LabID) Event and Infection Surveillance that may be tailored to meet the needs of participating NHSN facilities.

In addition, the following process measures are available: (1) adherence to hand hygiene; (2) adherence to contact precautions when caring for patients infected or colonized with an MDRO or *C. difficile*; and (3) adherence to active surveillance testing (AST) of MRSA and/or VRE. Active surveillance testing outcome measures is also available in locations where AST adherence is being performed and enables facilities to use the results of AST to monitor the incidence and prevalence of positive MRSA and/or VRE cultures. See the [MDRO/CDI](#) protocol for detailed surveillance instructions.

**Coronavirus Infectious Disease 2019 (COVID-19) Module**

The NHSN Coronavirus Infectious Disease 2019 Module was created during the COVID-19 outbreak as a means to monitor associated impacts on the human and material resources of acute care facilities. The module allows a facility to report information on the hospital’s capacity to respond to COVID-19 by reporting bed and mechanical ventilator availability as well as current and impending staffing and personal protective equipment shortages.
Outpatient Procedure Component Monthly Reporting Plan and Annual Facility Survey

Monthly Reporting Plan

The Outpatient Procedure Component (OPC) Monthly Reporting Plan form (CDC 57.401) is used by NHSN facilities to inform CDC which OPC modules are used during a given month. This allows CDC to select the data that should be included in the aggregate data analysis used for creating national benchmarks. Data entered into NHSN may represent either “in-plan” or “off-plan” surveillance. Each participating facility must identify and enter a monthly plan to indicate the module(s) used, if any, and the events, locations and/or procedures that will be monitored in-plan. The modules and locations selected for the month represent in-plan surveillance and indicate that the NHSN surveillance protocols will be used in their entirety for that surveillance.

- Only in-plan data are included in NHSN annual reports or other NHSN publications.
- “Off-plan” surveillance is surveillance that is done because a facility has decided to track a particular event for internal use. A facility makes no commitment to follow the NHSN protocol for “off-plan” events and such data are not included NHSN annual reports or other NHSN publications.

There must be a plan completed for every month that data are entered into NHSN although a facility may choose “No NHSN Outpatient Reporting this month” as an option. The reporting plan should take into account reporting requirements (for example, local or state mandates) when applicable to the facility.

Table of instructions for completing the Outpatient Procedure Component Monthly Reporting Plan form can be found at Table of Instructions for Completion of the Outpatient Procedure Component (OPC) Monthly Reporting Plan Form (CDC 57.401).

Ambulatory Surgery Center (ASC) Annual Facility Survey

The Outpatient Procedure Component (OPC) Annual Facility Survey (CDC 57.400) is used by CDC to classify facilities for appropriate comparisons in aggregate data analyses and to learn more about common practices among ASCs. Participating facilities must complete the Annual Facility Survey at the time that they enroll or activate the OPC and at the beginning of each calendar year thereafter. Most survey questions are based on facility characteristics and practices during the previous calendar year. For example, if a facility is completing the 2020 annual survey, they will report information that was recorded during January 2020 through December 2020.
Newly enrolled facilities that indicate that they were non-operational in the previous calendar year will be prompted to complete only a shortened version of the survey containing basic facility characteristic questions. At the start of the next calendar that follows when a facility enrolled into NHSN, they will complete the same survey, but will have to complete all the questions. For example, if the facility is enrolling in NHSN for the first time in March of 2020, they will complete a shortened version of the 2020 survey during enrollment. In January 2021, they will complete the entire 2020 survey with data from when they enrolled (March 2020) through December 2020.

Surveys must be completed by March 1st each year. After March 1st, facilities will be prevented from entering new monthly reporting plans until completion of the applicable survey(s).

The NHSN recommends that users collect all survey information using the paper form before attempting to enter data into the web application, as the application will not save incomplete surveys, meaning all required questions must be answered in order to complete and save the survey.

Instructions for completing the Annual Survey for Ambulatory Surgery Centers can be found at Table of Instructions for Completion of the Outpatient Procedure Component Annual Ambulatory Surgery Center Survey (CDC 57.400).
Same Day Outcome Measures (SDOM)

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Introduction

The measures that are included in this grouping of care quality indicators represent a broad range of risks encountered by patients accessing care in various outpatient settings. The four measures reflect the potential outcome resulting from procedures performed in the Ambulatory Surgery Center (ASC) outpatient environment. These potential outcomes can occur on the same day (during or immediately following) a procedure performed in an ASC. Same Day Outcome Measures includes four individual measures, which are:

1. Patient Burn
2. Patient Fall
3. Wrong Site, Wrong Side, Wrong Patient, Wrong Procedure, Wrong Implant
4. All-Cause Hospital Transfer/Admission

This NHSN protocol is intended to be consistent with the measure specifications from the following:

Key Terms for SDOM

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<td>Burn</td>
<td>Unintended tissue injury caused by any of the six recognized mechanisms: scalds, contact, fire, chemical, electrical or radiation, (for example, warming devices, prep solutions, electrosurgical unit or laser).</td>
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<td>Discharge</td>
<td>Occurs when the patient leaves the confines of the ASC.</td>
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<tr>
<td>Encounter</td>
<td>Any patient visit to an ASC where the patient completes the registration process upon entry into the facility. Some ASCs may refer to this as an admission into the facility.</td>
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<tr>
<td>Fall</td>
<td>A sudden, uncontrolled, unintentional, downward displacement of the body to the ground or other object, excluding falls resulting from violent blows or other purposeful actions. (National Center for Patient Safety).</td>
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<tr>
<td>Hospital transfer/admission</td>
<td>Any transfer/admission from an ASC directly to an acute care hospital including hospital emergency room.</td>
</tr>
<tr>
<td>Wrong</td>
<td>Procedure is performed in a way that is not consistent with what is documented in the informed consent for a patient - not in accordance with intended site, side, patient, procedure or implant</td>
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SDOM Requirements

Setting(s)
Any ASC as defined in the Code of Federal Regulations 42 CFR § 416.2 and has a “C” as the 3rd digit of its CMS Certification Number (CCN) eligible to use this measure.

Surveillance for SDOM
Monitor all patient encounters for the following events:
- Burn prior to discharge from the ASC
- Fall within the ASC
- Wrong site, side, patient, procedure or implant while admitted to the ASC (Wrong)
- Transfer or admission to a hospital upon discharge from the ASC

SDOM surveillance should be indicated on the NHSN Outpatient Procedure Component Monthly Reporting Plan form (CDC 57.401).

Monitoring for SDOM require active, patient-based, prospective surveillance. Surveillance for SDOMs starts at the beginning of the encounter and ends at discharge from the ASC. No post-discharge surveillance is required for these measures.
Methods for surveillance may vary based on resources within the facility. Examples of resources for data collection include outpatient facility medical records, incident/occurrence reports, or variance reports.

**Reporting these measures using NHSN SDOM does not meet the reporting requirement(s) for the CMS Ambulatory Surgical Center Quality Reporting (ASCQR) Program.** Reporting this NHSN measure is optional.

**SDOM Specifications**

➢ **Patient Burn**

There are several accounts in literature of patient burns in the surgical and procedural environment. The wide range of factors resulting in burns highlights several possible risks that must be addressed to prevent patient burns.

Many instances of burns are associated with electrosurgical equipment suggest that this is the most common causative agent. Recent reports demonstrate increased risk of burns may be related to newer devices that use higher currents at longer activation times. Although electrical burns may be the most predominant, burns from other mechanisms such as chemicals and direct contact have been reported.

Surgical fires are infrequent, but they are life threatening and the outcome (such as burns) can be severe to both patient and surgical staff. Any area where surgery is performed and flammable agents are used, such as medical gases and skin preparation agents, may pose a risk for surgical fires and subsequent patient burns.

Understanding that there are a number of causative agents related to patient burns in a surgical setting including ASCs, the term burn is very broad. This term covers burn from the various means by which a burn can occur – chemical, contact, electrical, fire radiation or scalds. This allows stakeholders and partners to gain a more robust understanding of the incidence of burn events and further improve prevention strategies.

**Measure Specifications:**

This measure is used to assess the number of encounters (patients) who experience a burn prior to discharge from the ASC.

- **Numerator:** ASC encounters (admissions) experiencing a burn prior to discharge
- **Exclusions:** None

- **Denominator:** All ASC encounters (admissions)
- **Exclusions:** None
➢ **Patient Fall**

The incidence of patient falls is currently unavailable, although in general the incidence of adverse events in ASCs is relatively low. There is growing interest in public reporting of adverse events such as falls. Patients undergoing outpatient surgical procedures are at increased risk for falls when adjunct therapies such as anxiolytics, sedatives, and anesthetic agents are used.

**Measure Specifications:**

This measure is used to assess the number of encounters (patients) who experience a fall within the ASC.

- **Numerator:** ASC encounters (admissions) experiencing a fall within the confines of the ASC
- **Exclusions:** ASC encounters (admissions) experiencing a fall outside the ASC

- **Denominator:** All ASC encounters (admissions)
- **Exclusions:** Falls resulting from violent blows or other purposeful actions

➢ **Wrong Site, Wrong Side, Wrong Patient, Wrong Procedure, and Wrong Implant** *(collectively referred to as Wrong)*

The Wrong outcome measure serves as a proxy for adherence to The Joint Commission’s “Universal Protocol” guideline. The goal for of the “Universal Protocol” guideline is to eliminate wrong site, wrong procedure, and wrong person surgery\(^1,2\). The “Universal Protocol” is a consensus guideline that is endorsed by professional medical organizations and associations. The ASC Quality Collaboration added wrong implant to wrong site, wrong side, wrong patient, and wrong procedure to create a more complete “wrong” event measure.

**Measure Specifications:**

This measure is used to assess the number of encounters (patients) who experience a wrong

- **Numerator:** All ASC encounters (admissions) experiencing a wrong
- **Exclusions:** None

- **Denominator:** All ASC encounters (admissions)
- **Exclusions:** None
➢ All-Cause Hospital Transfer/Admission

An unanticipated outcome after care is provided in an ASC, is a direct transfer or admission to a hospital from the ASCs. This unexpected event may result in additional cost and recovery time, which may pose an increased burden to the patient, family and payer.

At times, unforeseen events or complications may result in the need to transfer or admit an ASC patient to a hospital. Such occurrences demonstrate good judgement and signifies good patient care, but higher rates may be a signal that less than optimal patient and/or procedure selection by the ASC are occurring.

Measure Specifications:
This measure is used to assess the percentage of ASC encounters (admissions) who are transferred or admitted to a hospital upon discharge from the ASC

**Numerator:** ASC encounters (admissions) requiring a hospital transfer or hospital admission upon discharge from the ASC

**Exclusions:** None

**Denominator:** All ASC encounters (admissions)

**Exclusions:** None

Reporting Instructions

1. Indicate on the Outpatient Procedure Component Monthly Reporting Plan form (CDC 57.401) that the ASC is participating in surveillance for the Same Day Outcome Measures. Selecting SDOM means all four outcome measures will be monitored and reported.

2. For each patient that experiences a SDOM event during an ASC encounter, complete an Outpatient Procedure Component Same Day Outcome Measures Event form (CDC 57.402) and select the appropriate event by checking the corresponding box.
   a. If the same patient experiences more than one event of a different measure during the same encounter, all events should be recorded on the same event form. Example: a patient experiences a fall and a burn during the same encounter.
   b. If a patient experiences more than one event of the same measure during the same encounter, record only one event of that measure type for the encounter. Example: a patient has multiple wrong site procedures or multiple falls.
   c. If no events occur during an encounter, no Outpatient Procedure Component Same Day Outcome Measures Event form (CDC 57.402) should be completed

3. If no events occur during the reporting month, select “No Same Day Outcome Measures (events) reported this month” on the Outpatient Procedure Component Denominator for Same Day Outcome Measures form (CDC 57.403).
4. At the end of the reporting month specified in the Monthly Reporting Plan, enter the total number of ASC encounters (admissions) on the Outpatient Procedure Component Denominator for Same Day Outcome Measures form (CDC 57.403).

See the following for assistance with completing forms for the OPC Same Day Outcome Measures:

- *Table of Instructions for Completion of the Outpatient Procedure Component Monthly Reporting Plan Form (CDC 57.401)*
- *Table of Instructions for Completion of Outpatient Procedure Component Same Day Outcome Measures Form (CDC 57.402)*
- *Table of Instructions for Completion of the Outpatient Procedure Component Denominator for Same Day Outcome Measures Form (CDC 57.403)*

**Data Analysis**

Descriptive analysis options of numerator and denominator data, such as line listings, frequency tables, and bar and pie charts, are available in the NHSN application for analysis of SDOMs. Guides on using the NHSN OPC analysis features are available in the Analysis Resources section found at [https://www.cdc.gov/nhsn/ambulatory-surgery/ssi/index.html](https://www.cdc.gov/nhsn/ambulatory-surgery/ssi/index.html).
References


Outpatient Procedure Component Surgical Site Infection (OPC-SSI) Surveillance

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Introduction

With advances in surgical technology, patients are offered an incredible opportunity for restored health and function. The opposing force to technology advancements is increased risks of adverse and unintended outcomes such as surgical site infection (SSI). The CDC healthcare-associated infection (HAI) prevalence survey found that there were an estimated 110,800 surgical site infections (SSIs) associated with inpatient surgeries in 2015. As these data demonstrate, the frequency of SSI is primarily based on the analysis of operative procedures performed in inpatient settings such as acute care hospitals. These data represent only a fraction of the operative procedures performed on an annual basis and does not reflect the continued trend of surgical services transitioning to the outpatient ambulatory surgery settings.

In 2018, over 5717 Medicare-certified ambulatory surgery centers (ASCs), which represents nearly 17,400 operating rooms (ORs). This volume represents an average of 3.0 ORs per facility and an approximate 2.6-percent increase between 2017 and 2018. Therefore, it may be safe to assume that the continued growth in outpatient ORs equate to an increase in the volume of surgical procedures performed in the outpatient ambulatory surgery arena. Procedures performed in ambulatory surgery centers may be considered lower risk and thereby have a lower SSI rate than inpatient surgery settings, the continued growth in these facilities is a signal for the need to monitor procedures performed in the outpatient setting for adverse events such as SSIs. The OPC-SSI module will provide data for analyses to determine how operative procedures performed in ASCs contribute to the burden of SSIs. Data from this module can help identify factors associated with infections as well as targets for prevention strategies.

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. Surveillance of SSI with feedback of appropriate data to surgeons has been shown to be an important component of strategies to reduce SSI risk.

Advances have been made in infection control practices, including improved operating room ventilation, sterilization methods, barriers, surgical technique, and availability of antimicrobial prophylaxis, yet SSIs remain a substantial cause of morbidity, prolonged hospitalization, and death. Continued efforts are needed to identify preventable causes and develop strategies for SSI prevention.

The Outpatient Procedure Component (OPC) is designed for use by ambulatory surgery centers (ASCs). Surveillance for operative procedure(s) may focus on high risk and/or high-volume procedures. In addition, ASCs should use sound risk assessment practices as well as considerations for mandated reporting requirements to determine which operative procedure(s) to monitor. ASCs may voluntarily enroll in OPC-SSI but federal, state or organizational mandates supersedes voluntary enrollment and individual ASCs must verify and comply with mandated SSI reporting requirements.
OPC-SSI Reporting Requirements

OPC SSI reporting is based on the NHSN operative procedure categories. The NHSN operative procedure categories are listings of operative procedures grouped and categorized around a specific operative description. The operative procedure categories that are included in OPC-SSI surveillance can be found in Table 1. The Current Procedural Terminology (CPT) operative procedure codes are listed with the accompanying operative procedure code descriptions at https://www.cdc.gov/nhsn/xls/opc/opc-cpt-pcm-nhsn.xlsx.

Setting(s)

Any ASC as defined in the Code of Federal Regulations 42 CFR § 416.2 and has a “C” as the 3rd digit of its CMS Certification Number (CCN) is eligible to join NHSN OPC. These ASCs will use this protocol for surveillance of surgical patients receiving an eligible NHSN outpatient procedure (Table 1).

Reporting Plan

A facility may choose to perform surgical site surveillance “in-plan” or “off-plan” for any of the NHSN operative procedure categories:

- In-plan surveillance – Facility has indicated in their OPC Monthly Reporting Plan (CDC 57.401) that the OPC-SSI protocol will be used, in its entirety for SSI surveillance. Only in-plan data are included in NHSN annual reports or other NHSN publications.
- Off-plan surveillance – Facility has not indicated in their OPC Monthly Reporting Plan (CDC 57.401) that the OPC-SSI protocol will be used, in its entirety for SSI surveillance. Off-plan data are not included in NHSN annual reports or other NHSN publications.

Targeted Surveillance for OPC-SSI

a) For each calendar month under surveillance, indicate in the OPC Monthly Reporting Plan the NHSN operative procedure category in Table 1 that is under surveillance for SSI.

b) A facility may choose to monitor any of the NHSN operative procedure categories that are found in Table 1.

c) Perform surveillance for SSI following at least one NHSN operative procedure category (CPT Mapping) as indicated in the OPC Monthly Reporting Plan (CDC 57.401) and otherwise specified by mandates and other reporting requirements.

d) Collect SSI event (numerator) and operative procedure (denominator) data on all procedures included in the selected procedure category.

e) A procedure must meet the NHSN definition of an operative procedure in order to be included in the surveillance. All procedures included in the NHSN monthly surveillance plan are followed for all SSI types: superficial incisional, deep incisional, and organ/space. The type of SSI reported must reflect the deepest tissue level (superficial, deep and organ/space) where SSI criteria are met during the surveillance period.

NOTES:
• An SSI event is attributed to the facility in which the NHSN operative procedure was originally performed.
• Facilities that have identified potential SSI events that are attributable to procedures performed at a different facility should provide details of the potential events to the facility where the procedure was originally performed.

Table 1. NHSN OPC Operative Procedure Categories

<table>
<thead>
<tr>
<th>Procedure Category</th>
<th>Operative Procedure</th>
<th>Procedure Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>AMP</td>
<td>Limb amputation</td>
<td>Total or partial amputation or disarticulation of the upper or lower limbs, including digits</td>
</tr>
<tr>
<td>APPY</td>
<td>Appendix surgery</td>
<td>Operation of appendix</td>
</tr>
<tr>
<td>AVSD</td>
<td>AV shunt for dialysis</td>
<td>Arteriovenostomy for renal dialysis</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 on gall bladder only)</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>
</tr>
<tr>
<td>CEA</td>
<td>Carotid endarterectomy</td>
<td>Endarterectomy on vessels of head and neck (includes carotid artery and jugular vein)</td>
</tr>
<tr>
<td>CHOL</td>
<td>Gallbladder surgery</td>
<td>Cholecystectomy and cholecystotomy</td>
</tr>
<tr>
<td>COLO</td>
<td>Colon surgery</td>
<td>Incision, resection, or anastomosis of the large intestine; includes large-to-small and small-to-large bowel anastomosis; see REC for rectal operations</td>
</tr>
<tr>
<td>FUSN</td>
<td>Spinal fusion</td>
<td>Immobilization of spinal column</td>
</tr>
<tr>
<td>FX</td>
<td>Open reduction of fracture</td>
<td>Open reduction of fracture or dislocation of long bones without internal or external fixation; does not include placement of joint prosthesis</td>
</tr>
<tr>
<td>GAST</td>
<td>Gastric surgery</td>
<td>Incision or excision of stomach; includes subtotal or total gastrectomy; does not include vagotomy and fundoplication</td>
</tr>
<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>
</tr>
<tr>
<td>HPRO</td>
<td>Hip prosthesis</td>
<td>Arthroplasty of hip</td>
</tr>
<tr>
<td>HYST</td>
<td>Abdominal hysterectomy</td>
<td>Abdominal hysterectomy; includes that by laparoscope</td>
</tr>
<tr>
<td>KPRO</td>
<td>Knee prosthesis</td>
<td>Arthroplasty of knee</td>
</tr>
<tr>
<td>LAM</td>
<td>Laminectomy</td>
<td>Exploration or decompression of spinal cord through excision or incision into vertebral structures</td>
</tr>
</tbody>
</table>
### NHSN Operative Procedure Category Mappings to CPT Codes

Operative procedure codes are used in various health care settings as a uniform way to communicate essential information. This wide use of operative procedure codes allows NHSN to standardize the SSI surveillance reporting process. **Current Procedural Terminology (CPT) codes are the operative procedure codes used in OPC and are required for use within the application.**

NHSN has mapped Current Procedural Terminology (CPT) codes to NHSN OPC operative procedure categories to assist in determining the correct operative procedures to report for SSI surveillance. The CPT mapping to OPC operative procedure categories can be found in the “Supporting Materials” section of the ASC SSI Events webpage. The CPT mapping document includes a general definition for each OPC operative procedure category as well as a procedure description for each individual operative procedure code.

<table>
<thead>
<tr>
<th>Category</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>NECK</td>
<td>Neck surgery Major excision or incision of the larynx and radical neck dissection; does not include thyroid and parathyroid operations</td>
</tr>
<tr>
<td>NEPH</td>
<td>Kidney surgery Resection or manipulation of the kidney with or without removal of related structures</td>
</tr>
<tr>
<td>OVRY</td>
<td>Ovarian surgery Operations on ovary and related structures</td>
</tr>
<tr>
<td>PACE</td>
<td>Pacemaker surgery Insertion, manipulation or replacement of pacemaker</td>
</tr>
<tr>
<td>PRST</td>
<td>Prostate surgery Suprapubic, retropubic, radical, or perineal excision of the prostate; does not include transurethral resection of the prostate</td>
</tr>
<tr>
<td>PVBY</td>
<td>Peripheral vascular bypass surgery Bypass operations on peripheral arteries and veins</td>
</tr>
<tr>
<td>REC</td>
<td>Rectal surgery Operations on rectum</td>
</tr>
<tr>
<td>SB</td>
<td>Small bowel surgery Incision or resection of the small intestine; does not include small-to-large bowel anastomosis</td>
</tr>
<tr>
<td>SPLE</td>
<td>Spleen surgery Resection or manipulation of spleen</td>
</tr>
<tr>
<td>THOR</td>
<td>Thoracic surgery Noncardiac, nonvascular thoracic surgery; includes pneumonectomy and hiatal hernia repair or diaphragmatic hernia repair (except through abdominal approach)</td>
</tr>
<tr>
<td>THYR</td>
<td>Thyroid and/or parathyroid surgery Resection or manipulation of thyroid and/or parathyroid</td>
</tr>
<tr>
<td>VHYS</td>
<td>Vaginal hysterectomy Vaginal hysterectomy; excludes the use of laparoscope</td>
</tr>
<tr>
<td>VSHN</td>
<td>Ventricular shunt Ventricular shunt operations, including revision and removal of shunt</td>
</tr>
<tr>
<td>XLAP</td>
<td>Exploratory laparotomy Abdominal operations not involving the gastrointestinal tract or biliary system; includes diaphragmatic hernia repair through abdominal approach</td>
</tr>
</tbody>
</table>
Custom Procedures, Custom Events and Custom Fields

Custom procedures, custom events and custom fields are created by individual facilities. These custom data are optional and allow facility-defined data entry for the facility’s own surveillance purposes.

- Custom procedures are non-NHSN operative procedures and cannot be included in the Monthly Reporting Plan and are therefore considered off-plan surveillance.
- Custom events are non-NHSN defined events based on criteria developed by the facility.
- Custom fields are non-NHSN defined variables. These fields may be added to NHSN-defined procedures.

Custom fields, custom procedures and custom events must be created in the application before data can be entered. These data may provide value to the facility if they are entered in a consistent manner. For example, if a facility chooses to create a custom field for admission or discharge diagnosis of infected patients, standardized responses should be entered in a consistent and uniform manner.

Data entered in custom fields or in association with custom procedures and events are not included in any of the NHSN reports and there are no available NHSN comparative data. Any related analyses must be performed by the facility.

Instructions for creating custom fields, procedures and events may be found in the Supporting Materials section of the Ambulatory Surgery Center webpage.

Key Terms for OPC-SSI

**Physician** - should be interpreted to mean the surgeon(s), infectious disease physician, emergency physician, other physician on the case, or physician's designee (nurse practitioner or physician's assistant).

**Date of event (DOE)** - the date when the first element used to meet the OPC-SSI infection criterion occurs for the first time during the SSI surveillance period. The date of event must fall within the SSI surveillance period to meet SSI criteria. The type of SSI (superficial incisional, deep incisional, or organ/space) reported should reflect the deepest tissue layer involved in the infection during the surveillance period. Synonym: infection date.

**NOTE:**
All elements (signs/symptoms) required to meet an SSI criterion usually occur within a 7-10-day timeframe with no more than 2-3-days between elements. The elements must be relational to each other, meaning all elements associated with the SSI occur in a relatively tight timeframe. Each case differs based on the individual criteria elements occurring and the type of SSI.

**NHSN Operative Procedure** - is a procedure that
• is included in the NHSN CPT operative procedure category code mapping
  and
• takes place during an operation where at least one incision (including laparoscopic approach) is
  made through the skin or mucous membrane, or reoperation via an incision that was left open
  during a prior operative procedure
  and
• takes place in an operating room (OR), defined as a patient care area that met criteria for an
  operating room when it was constructed or renovated outlined by the Facilities Guidelines
  Institute’s (FGI)°, American Institute of Architects’ (AIA) or requirements of the State in which it
  operates. This may include an interventional radiology room, or a cardiac catheterization lab.

**Surveillance Period** - is the timeframe following an NHSN operative procedure for monitoring and
identifying a post-operative infection, see Table 2. The surveillance period is determined by the NHSN
operative procedure category (for example, BRST-Breast surgery has a 90-day surveillance period and
HYST-abdominal hysterectomy surgeries have a 30-day surveillance period).
Table 2. Surveillance Periods for SSIs Following Selected NHSN Operative Procedure Categories. Day 1 = the date of the procedure.

<table>
<thead>
<tr>
<th>30-day Surveillance</th>
<th>Category</th>
<th>Operative Procedure</th>
<th>Category</th>
<th>Operative Procedure</th>
</tr>
</thead>
<tbody>
<tr>
<td>AMP</td>
<td>Limb amputation</td>
<td>NECK</td>
<td>Neck surgery</td>
<td></td>
</tr>
<tr>
<td>APPY</td>
<td>Appendix surgery</td>
<td>NEPH</td>
<td>Kidney surgery</td>
<td></td>
</tr>
<tr>
<td>AVSD</td>
<td>Shunt for dialysis</td>
<td>OVRY</td>
<td>Ovarian surgery</td>
<td></td>
</tr>
<tr>
<td>BILI</td>
<td>Bile duct, liver or pancreatic surgery</td>
<td>PRST</td>
<td>Prostate surgery</td>
<td></td>
</tr>
<tr>
<td>CEA</td>
<td>Carotid endarterectomy</td>
<td>REC</td>
<td>Rectal surgery</td>
<td></td>
</tr>
<tr>
<td>CHOL</td>
<td>Gallbladder surgery</td>
<td>SB</td>
<td>Small bowel surgery</td>
<td></td>
</tr>
<tr>
<td>COLO</td>
<td>Colon surgery</td>
<td>SPLE</td>
<td>Spleen surgery</td>
<td></td>
</tr>
<tr>
<td>GAST</td>
<td>Gastric surgery</td>
<td>THOR</td>
<td>Thoracic surgery</td>
<td></td>
</tr>
<tr>
<td>HYST</td>
<td>Abdominal hysterectomy</td>
<td>THYR</td>
<td>Thyroid and/or parathyroid surgery</td>
<td></td>
</tr>
<tr>
<td>LAM</td>
<td>Laminectomy</td>
<td>VHYS</td>
<td>Vaginal hysterectomy</td>
<td></td>
</tr>
<tr>
<td>-</td>
<td>-</td>
<td>XLAP</td>
<td>Exploratory Laparotomy</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>90-day Surveillance</th>
<th>Category</th>
<th>Operative Procedure</th>
</tr>
</thead>
<tbody>
<tr>
<td>BRST</td>
<td>Breast surgery</td>
<td></td>
</tr>
<tr>
<td>FUSN</td>
<td>Spinal fusion</td>
<td></td>
</tr>
<tr>
<td>FX</td>
<td>Open reduction of fracture</td>
<td></td>
</tr>
<tr>
<td>HER</td>
<td>Herniorrhaphy</td>
<td></td>
</tr>
<tr>
<td>HPRO</td>
<td>Hip prosthesis</td>
<td></td>
</tr>
<tr>
<td>KPRO</td>
<td>Knee prosthesis</td>
<td></td>
</tr>
<tr>
<td>PACE</td>
<td>Pacemaker surgery</td>
<td></td>
</tr>
<tr>
<td>PVBY</td>
<td>Peripheral vascular bypass surgery</td>
<td></td>
</tr>
<tr>
<td>VSHN</td>
<td>Ventricular shunt</td>
<td></td>
</tr>
</tbody>
</table>

**NOTES:**
- Superficial incisional SSIs are only followed for a 30-day period for all procedure types.
- Secondary incisional SSIs are only followed for a 30-day period regardless of the surveillance period for the primary site.
Table 3. Denominator for Procedure Details

These are required elements for reporting each operative procedure performed within the selected operative procedure category. The elements have been identified as risk factors for SSIs. See the *Instructions for Completion of Outpatient Procedure Component (OPC) Denominator for Procedure Form (CDC 57.404)* for further details.

<table>
<thead>
<tr>
<th>Element Description</th>
<th>Description</th>
</tr>
</thead>
</table>
| ASA physical status | Assessment by the anesthesiologist of the patient’s preoperative physical condition using the American Society of Anesthesiologists’ (ASA) Physical Status Classification System\(^1\). At the time of surgery, the patient is assigned an ASA score of 1-6.  
**NOTES:**  
- *Do NOT report procedures that do not have an ASA score assigned by an anesthesiologist.*  
- *Do NOT report procedures with an ASA physical status of 6 (a declared brain-dead patient whose organs are being removed for donor purposes) to NHSN.* |
| Diabetes | The NHSN SSI surveillance definition of diabetes indicates that the patient has a diagnosis of diabetes requiring management with insulin or a non-insulin anti-diabetic agent. This include patients with:  
- “Insulin resistance” who are on management with anti-diabetic agents.  
- A diagnosis of diabetes who are noncompliant with their diabetes medications.  
- Gestational diabetes.  
ICD-10-CM diagnosis codes (that reflect a diagnosis of diabetes) documented during the admission when the procedure is performed maybe used to determine diabetes. Acceptable codes are found in the “Supporting Materials” section of the ASC SSI Events webpage.  
**NOTE:**  
The NHSN definition excludes patients with no diagnosis of diabetes. The definition also excludes patients who receive insulin only for perioperative control of hyperglycemia but have no diagnosis of diabetes. |
| Duration of operative procedure | The interval in hours and minutes between the Procedure/Surgery Start Time, and the Procedure/Surgery Finish Time, as defined by the Association of Anesthesia Clinical Directors (AACD)\(^5\):  
- Procedure/Surgery Start Time (PST): Time when the procedure is begun (for example, incision for a surgical procedure).  
- Procedure/Surgery Finish (PF): Time when all instrument and sponge counts are completed and verified as correct, all postoperative radiologic studies to be done in the OR are completed, all dressings and drains are secured, and the physicians/surgeons have completed all procedure-related activities on the patient. |
| **General anesthesia** | The administration of drugs or gases that enter the general circulation and affect the central nervous system to render the patient pain free, amnesic, unconscious, and often paralyzed with relaxed muscles. This does not include conscious sedation. |
| **Height** | The patient’s most recent height documented in the medical record in feet (ft.) and inches (in.) or meters (m). |
| **Scope** | An instrument used to visualize the interior of a body cavity or organ. In the context of an NHSN operative procedure, use of a scope involves creation of several small incisions to perform or assist in the performance of an operation rather than use of a traditional larger incision (specifically, open approach).

For CPT codes, the scope question can be answered based on the procedure code description. Using HYST code 58570 as an example, the procedure code description indicates Laparoscopy, surgical, with total hysterectomy. Laparoscopy is **Scope = YES**.

| HYST | 58570 | Laparoscopy, surgical, with total hysterectomy, for uterus 250 g or less |
| **Weight** | The patient’s most recent weight documented in the medical record in pounds (lbs.) or kilograms (kg) prior to or otherwise closest to the procedure. |
| **Wound class** | An assessment of the degree of contamination of a surgical wound at the time of the operation. Wound class should be assigned by a person involved in the surgical procedure *(for example, surgeon, circulating nurse, etc.)*. The four wound classes available within the NHSN application are Clean, Clean-Contaminated, Contaminated, and Dirty/Infected. Each facility should use the wound class schema as stated in its policies and procedures for clinical practices.

**NOTE:**
Based on feedback from external experts in the field of surgery, NHSN surgical procedure categories APPY - Appendix surgery, BILI - Bile duct, liver or pancreatic surgery, CHOL - Gallbladder surgery, COLO - Colon surgery, REC - Rectal surgery, SB - Small bowel surgery and VHYS - Vaginal hysterectomy should never be recorded as clean wound class. The rationale for this is due to the anatomy of the body and the usual approach required to reach the operative site. For these operative procedures clean wound class is not an option on the drop-down menu within the application.
**NOTE:**

*Incisional closure method is NOT a part of the NHSN OPC-SSI Surveillance definition; therefore, all eligible procedures should be included in SSI surveillance regardless of closure method. Both primarily closed procedures and those that are not closed primarily should be included in the denominator data for procedures in the facility’s NHSN Monthly Reporting Plan. Any SSI attributable to either primarily closed or non-primarily closed procedures should be reported.*
Surgical Site Infection (SSI) Criteria

Table 4A: General OPC-SSI Criteria

**APPLY TO ALL OPERATIVE PROCEDURE CATEGORIES EXCEPT BREAST SURGERY (BRST). USE BREAST SURGERY (BRST) - SURGICAL SITE INFECTION CRITERIA FOR SSIS ATTRIBUTABLE TO BRST.**

<table>
<thead>
<tr>
<th>OPC General – Superficial Incisional SSI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Must meet the following criteria:</td>
</tr>
<tr>
<td>Date of event for infection occurs within 30 days after any NHSN operative procedure (where day 1 = the procedure date)</td>
</tr>
<tr>
<td><strong>AND</strong></td>
</tr>
<tr>
<td>involves only skin and subcutaneous tissue of the incision</td>
</tr>
<tr>
<td><strong>AND</strong></td>
</tr>
<tr>
<td>patient has at least one of the following:</td>
</tr>
<tr>
<td>a. purulent drainage from the superficial incision.</td>
</tr>
<tr>
<td>b. organisms identified from an aseptically-obtained specimen from the superficial incision or subcutaneous tissue by a culture or non-culture based microbiologic testing method which is performed for purposes of clinical diagnosis or treatment (for example, not Active Surveillance Culture/Testing [ASC/AST]).</td>
</tr>
<tr>
<td>c. superficial incision that is deliberately opened by a surgeon, physician or physician designee and culture or non-culture based testing of the superficial incision or subcutaneous tissue is not performed.</td>
</tr>
<tr>
<td><strong>and</strong></td>
</tr>
<tr>
<td>patient has at least one of the following signs or symptoms: localized pain or tenderness; localized swelling; erythema; or heat.</td>
</tr>
<tr>
<td>d. diagnosis of a superficial incisional SSI by a physician or physician designee.</td>
</tr>
</tbody>
</table>

**Comments:** The two specific types of superficial incisional SSIs are:

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 (for example, the knee incision for KPRO procedure).

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 (for example, abdominal incision site for VSHN).
### Reporting Instructions for OPC General - Superficial Incisional SSI

The following do not qualify as criteria for meeting the NHSN definition of superficial SSI:

- Diagnosis/treatment of cellulitis (redness/warmth/swelling), by itself, does not meet criterion “d” for superficial incisional SSI. Conversely, an incision that is draining or that has organisms identified by culture or non-culture based testing is not considered a cellulitis.
- A stitch abscess alone (minimal inflammation and discharge confined to the points of suture penetration).
- A localized stab wound, or pin site infection is not an SSI.

**NOTE:**

*A laparoscopic trocar site for an NHSN operative procedure is not considered a stab wound.*

### OPC General - Deep Incisional SSI

Must meet the following criteria:

- The date of event for infection occurs within 30 or 90 days after the NHSN operative procedure (where day 1 = the procedure date) according to the list in Table 2
- **AND**
  - involves deep soft tissues of the incision (for example, fascial and muscle layers)
  - **AND**
    - patient has at least one of the following:
      - a. purulent drainage from the deep incision.
      - b. a deep incision that spontaneously dehisces, or is deliberately opened or aspirated by a surgeon, physician or physician designee.
        **and**
        - organism is identified from the deep soft tissues of the incision by a culture or non-culture based microbiologic testing method which is performed for purposes of clinical diagnosis or treatment (for example, not Active Surveillance Culture/Testing [ASC/AST]) or culture or non-culture based microbiologic testing method is not performed. A culture or non-culture based test from the deep soft tissues of the incision that has a negative finding does not meet this criterion.
        **and**
        - patient has at least one of the following signs or symptoms: fever (>38°C); localized pain or tenderness.
      - c. an abscess or other evidence of infection involving the deep incision that is detected on gross anatomical or histopathologic exam, or imaging test.
**Comments:** The two specific types of deep incisional SSIs are:

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 (for example, the hip incision for a HPRO procedure).

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 (for example, abdominal incision site for VSHN).

**OPC General - Organ/Space SSI**
Must meet the following criteria:

- Date of event for infection occurs within 30 or 90 days after the NHSN operative procedure (where day 1 = the procedure date) according to the list in Table 2
- AND
- infection involves any part of the body deeper than the fascial/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 into the organ/space (for example, closed suction drainage system, open drain, T-tube drain, and CT guided drainage).
  - b. organisms are identified from an aseptically-obtained fluid or tissue in the organ/space by a culture or non-culture based microbiologic testing method which is performed for purposes of clinical diagnosis or treatment (for example, not Active Surveillance Culture/Testing [ASC/AST]).
  - c. an abscess or other evidence of infection involving the organ/space that is detected on gross anatomical or histopathologic exam, or imaging test consistent with infection.

**NOTE:**
*Meeting additional infection criteria found in the Patient Safety Component Chapter 17, CDC/NHSN Surveillance Definitions for Specific Types of Infections is NOT a part of the OPC General - Organ/Space SSIs reporting criteria.*
Table 4B: Breast Surgery (BRST) Surgical Site Infection Criteria

The Breast Surgery (BRST) Surgical Site Infection instructions apply to surgical site infections (SSIs) during the 30-day (superficial SSI) and 90-day (deep and organ/space SSI) postoperative periods following BRST-Breast Surgery performed in Ambulatory Surgery Centers. Use General OPC-SSI criteria for all operative procedures except breast surgery (BRST).

<table>
<thead>
<tr>
<th>OPC BRST - Superficial incisional SSI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Must meet the following criteria:</td>
</tr>
<tr>
<td>Date of event for infection occurs within 30 days after a BRST; where day 1 = the procedure date AND</td>
</tr>
<tr>
<td>involves either the skin, subcutaneous tissue (for example, fatty tissue) or breast parenchyma (for example, milk ducts and glands that produce milk) at the incision AND</td>
</tr>
<tr>
<td>patient has at least one of the following:</td>
</tr>
<tr>
<td>a. purulent drainage from the superficial incision.</td>
</tr>
<tr>
<td>b. organisms identified from an aseptically-obtained specimen from the superficial incision or subcutaneous tissue by a culture or non-culture based microbiologic testing method which is performed for purposes of clinical diagnosis or treatment (for example, not Active Surveillance Culture/Testing [ASC/AST]).</td>
</tr>
<tr>
<td>c. superficial incision that is deliberately opened by a surgeon, physician or physician designee and culture or non-culture based testing of the superficial incision or subcutaneous tissue is not performed. and</td>
</tr>
<tr>
<td>patient has at least one of the following signs or symptoms: localized pain or tenderness; localized swelling; redness (erythema); or heat. A culture or non-culture based test that has a negative finding does not meet this criterion.</td>
</tr>
<tr>
<td>d. diagnosis of a superficial incisional SSI by a physician or physician designee.</td>
</tr>
</tbody>
</table>

Comments for OPC BRST – Superficial Incisional SSI

The two specific types of superficial incisional SSIs are:

1. Superficial incisional primary (SIP) – a superficial incisional SSI that is identified in a primary incision in a patient that has had an operation with one or more incisions (for example, the breast incision for BRST procedure).

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 (for example, transverse rectus abdominis myocutaneous [TRAM] flap incision site for BRST).
**OPC BRST - Deep incisional SSI**

Must meet the following criteria:

- Date of event for infection occurs within 90 days after a BRST; where day 1 = the procedure date
- **AND**
  - involves deep soft tissues of the incision (for example, fascial and muscle layers)
- **AND**
  - patient has at least one of the following:
    - a. purulent drainage from the deep incision.
    - b. a deep incision that spontaneously dehiscences, or is deliberately opened or aspirated by a surgeon, physician or physician designee.
      - **and**
      - organism is identified from the deep soft tissues of the incision by a culture or non-culture based microbiologic testing method which is performed for purposes of clinical diagnosis or treatment (for example, not Active Surveillance Culture/Testing [ASC/AST]) or culture or non-culture based microbiologic testing method is not performed. A culture or non-culture based test that has a negative finding does not meet this criterion.
      - **and**
      - patient has at least **one** of the following signs or symptoms: fever (>38°C); localized pain or tenderness.
    - c. an abscess or other evidence of infection involving the deep incision that is detected on gross anatomical or histopathologic exam.

**Comments for OPC BRST – Deep Incisional SSI**

The two specific types of deep incisional SSIs are:

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 (for example, the breast incision for BRST procedure).

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 (for example, transverse rectus abdominis myocutaneous [TRAM] flap incision site for BRST).
### OPC BRST - Organ/Space SSI

Must meet the following criteria:

**Date of event for infection occurs within 90 days a BRST; where day 1 = the procedure date**

**AND**

- infection involves any part of the body deeper than the fascial/muscle layers (subpectoral), that is opened or manipulated during the operative procedure.

**AND**

- patient has at least one of the following:
  1. purulent drainage from a drain that is placed into the organ/space (for example, closed suction drainage system, open drain, T-tube drain, and CT guided drainage).
  2. organisms identified from affected breast tissue or fluid obtained by invasive procedure by a culture or non-culture based microbiologic testing method which is performed for purposes of clinical diagnosis or treatment (for example, not Active Surveillance Culture/Testing [ASC/AST]).
  3. breast abscess or other evidence of infection on gross anatomic or histopathologic exam or imaging test consistent with breast infection.

### NOTES:

- **Breast surgeries may involve a secondary operative incision, specifically, procedures that include a transverse rectus abdominis flap. The flap site is the secondary operative incision. Secondary sites have a 30-day surveillance period. If the secondary site meets criteria for an SSI, it is reported as either a superficial incisional SSI at the secondary site or deep incisional infection at the incisional site.**
- **Accessing a breast expander after a breast surgery is considered an invasive procedure and any subsequent infection is not deemed an SSI attributable to the breast surgery.**
- **Meeting additional infection criteria found in the Patient Safety Component Chapter 17, CDC/NHSN Surveillance Definitions for Specific Types of Infections is NOT a part of the OPC BRST - Organ/Space SSIs reporting criteria.**
OPC-SSI Numerator (SSI Event) Reporting

Numerator Data

a) All patients having any of the procedures included in the selected NHSN operative procedure category(s) are monitored for SSI. The Outpatient Procedure Component (OPC) Surgical Site Infection (SSI) Event Form (CDC 57.405) is completed for each SSI.

b) If no SSI events are identified during the surveillance month, check the “Report No Events” field in the Missing OPC Events tab of the Incomplete/Missing List.

c) The Instructions for the Completion of Outpatient Procedure Component Surgical Site Infection (OPC-SSI) Event Form (CDC 57.405) form include brief instructions for collection and entry of each data element on the form. The OPC-SSI data collection form includes patient demographic information and information about the operative procedure, including the date and type of procedure. As well as information about the SSI including the date of SSI, specific criteria met for identifying the SSI and when/how the SSI was detected.

d) See the OPC tables of instructions for detailed information regarding the completion of the OPC Monthly Reporting Plan Form (CDC 57.401), Outpatient Procedure Component (OPC) Denominator for Procedure Form (CDC 57.404), and SSI information for the Outpatient Procedure Component (OPC) Surgical Site Infection (SSI) Event Form (CDC 57.405).

Table 5: Numerator Reporting Instructions

Numerator (SSI event) reporting instructions are guidelines for reporting SSI events. The instructions ensure consistent application of the general and breast surgery reporting criteria.

<table>
<thead>
<tr>
<th>Topic</th>
<th>Reporting Instruction</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Excluded organisms:</td>
<td>Well-known community associated (organisms belonging to the following genera: Blastomyces, Histoplasma, Coccidioides, Paracoccidioides, Cryptococcus and Pneumocystis and/or organisms associated with latent infections (for example, herpes, shingles, syphilis, or tuberculosis) are excluded from meeting SSI criteria.</td>
</tr>
<tr>
<td>2. Attributing SSI to an NHSN procedure when there is evidence of infection at the time of the primary surgery:</td>
<td>SSI surveillance does not take into account infections that are present at the operative site at the time of the operative procedure. When there is evidence of an infection at the operative site at the time of the operative procedure and if during the SSI surveillance period the patient meets NHSN OPC-SSI criteria, an SSI should be attributed to the operative procedure. A procedure with a high wound class is included in denominator reporting and is eligible for SSI surveillance; in many cases, wound class is included as a risk factor for SSI in the NHSN risk modeling.</td>
</tr>
</tbody>
</table>
### Reporting Instruction

#### 3. Multiple tissue levels are involved in the infection:

The type of SSI (superficial incisional, deep incisional, or organ/space) reported must reflect the deepest tissue level where SSI criteria are met during the surveillance period. The date of event should be the date that the patient met criteria for the deepest level of infection.

For example:

- Report infection that involves the organ/space as an organ/space SSI, whether or not it also involves the superficial or deep incision levels.
- Report infection that involves the superficial and deep incisional levels as a deep incisional SSI.
- If an SSI started as a deep incisional SSI on day 10 of the SSI surveillance period and then a week later, (day 17 of the SSI surveillance period) meets criteria for an organ space SSI the date of event would be the date of the organ space SSI.

#### 4. Attributing SSI to NHSN procedures that involve multiple primary incision sites:

If multiple primary incision sites of the same NHSN operative procedure become infected, only report as a single SSI, and assign the type of SSI (superficial incisional, deep incisional, or organ/space) that represents the deepest tissue level where SSI criteria is met at any of the infected involved primary incision sites during the surveillance period.

For example:

- If one laparoscopic incision meets criteria for a superficial incisional SSI and another meets criteria for a deep incisional SSI, only report one deep incisional SSI.
- If one or more laparoscopic incision sites meet criteria for superficial incisional SSI but the patient also has an organ/space SSI related to the laparoscopic procedure, only report one organ/space SSI.
- If an operative procedure is limited to a single breast and involves multiple incisions in that breast that become infected, only report a single SSI.
- In a colostomy formation or reversal (take down) procedure, the stoma and other abdominal incision sites are considered primary incisions. If both the stoma and another abdominal incision site develop superficial incisional SSI, report only as one SSI (SIP).
<table>
<thead>
<tr>
<th>Topic</th>
<th>Reporting Instruction</th>
</tr>
</thead>
</table>
| 5. Attributing SSI to NHSN procedures that have secondary incision sites: | Certain procedures can involve a secondary operative incision (for example, BRST, FUSN, PVBY, REC and VSHN). The surveillance period for all secondary operative incisions is 30 days, regardless of the required deep incisional or organ/space SSI surveillance period for the primary incision site(s) (Table 2). Procedures meeting this designation are reported as one (a single) operative procedure. For example:  
  - A tissue harvest site in a BRST procedure with a transverse rectus abdominis myocutaneous (TRAM) flap is considered the secondary operative incision. One BRST procedure is reported, and if the secondary incision becomes infected, report as either SIS or DIS as appropriate. |
| 6. SSI detected at another facility:                                   | It is required that if an SSI is detected at a facility other than the ASC where the procedure was originally performed, details of the SSI event should be provided to the ASC so the SSI can be accurately reported to NHSN. When reporting the SSI, the ASC should indicate how the SSI was identified/detected in the “SSI Event Detected” section of the OPC-SSI form. An SSI event is attributed to the facility in which the NHSN operative procedure was performed. For example:  
  - A patient had a fusion (FUSN) of the left sacroiliac joint preformed at an ASC. 35 days post-operative the patient was seen in the emergency department of a community hospital with signs and symptoms of infection at the surgical site. The community hospital contacted the ASC to report the patient’s signs and symptoms of infection at the left sacroiliac joint. Upon meeting OPC-SSI criteria the ASC should select, “Report from another facility (inpatient, health department, emergency department, etc.” in the SSI Event Detected” section of the OPC-SSI event form.  
  - An ASC has a formal post-discharge surveillance process which includes post-operative phone calls to the patient as well as surveys mailed to the surgeons. A surgeon returns a survey and notes a patient having had a breast surgery (BRST) was seen in his office with a superficial infection and was treated with an oral antibiotic. The ASC should select “Post-discharge surgeon survey” in the SSI Event Detected” section of the OPC-SSI event form. |
<p>| 7. SSI attribution after multiple types of NHSN                       | If more than one NHSN operative procedure category was performed through a single incision/laparoscopic sites during a |</p>
<table>
<thead>
<tr>
<th>Topic</th>
<th>Reporting Instruction</th>
</tr>
</thead>
<tbody>
<tr>
<td>procedures are performed during a single trip to the OR:</td>
<td>Single trip to the operating room, attribute the SSI to 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 an incisional SSI, use the NHSN Principal Operative Procedure Category Selection Lists (Table 6) to select the operative procedure to which the SSI should be attributed. For example: If a patient develops an SSI after a single trip to the OR in which both a COLO and SB were performed, and the source of the SSI is not apparent, assign the SSI to the COLO procedure.</td>
</tr>
<tr>
<td>8. SSI following invasive manipulation/accession of the operative site:</td>
<td>An SSI will NOT be attributed if ALL criteria are met (all three must be present): • during the post-operative period the surgical site is without evidence of infection and • an invasive manipulation/accession of the site is performed for diagnostic or therapeutic purposes (for example, needle aspiration, accession of ventricular shunts, accession of breast expanders) and • an infection subsequently develops in a tissue level which was entered during the manipulation/accession. Tissue levels that are BELOW the deepest level of manipulation/accession will be eligible for SSI. For example, in a superficial debridement following a COLO procedure, where the muscle/fascia and organ/space are not entered, a subsequent organ/space SSI following the debridement may be an SSI attributable to the index COLO procedure. This reporting instruction does NOT apply to closed manipulation (for example, closed reduction of a dislocated hip after an orthopedic procedure). Invasive manipulation does not include wound packing or changing of wound packing materials as part of postoperative care.</td>
</tr>
<tr>
<td>9. SSI following specific post-operative infection scenarios:</td>
<td>An SSI that otherwise meets the NHSN definitions should be reported to NHSN without regard to post-operative accidents, falls, inappropriate showering or bathing practices, or other occurrences that may or may not be attributable to patients’ intentional or unintentional postoperative actions.</td>
</tr>
</tbody>
</table>
SSI should also be reported regardless of the presence of certain skin conditions (for example, dermatitis, blister, impetigo) that occur near an incision, and regardless of the possible occurrence of a “seeding” event from an unrelated procedure (for example, dental work). These instructions concerning various postoperative circumstances are necessary to reduce subjectivity and data collection burden associated with the previously exempted scenarios.

Table 6. NHSN Principal Operative Procedure Category Selection List

(The categories with the highest risk of SSI are listed before those with lower risks).

<table>
<thead>
<tr>
<th>Priority</th>
<th>Procedure Category</th>
<th>Abdominal Operations</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>COLO</td>
<td>Colon surgery</td>
</tr>
<tr>
<td>2</td>
<td>BILI</td>
<td>Bile duct, liver or pancreatic surgery</td>
</tr>
<tr>
<td>3</td>
<td>SB</td>
<td>Small bowel surgery</td>
</tr>
<tr>
<td>4</td>
<td>REC</td>
<td>Rectal surgery</td>
</tr>
<tr>
<td>5</td>
<td>GAST</td>
<td>Gastric surgery</td>
</tr>
<tr>
<td>6</td>
<td>HYST</td>
<td>Abdominal hysterectomy</td>
</tr>
<tr>
<td>7</td>
<td>XLAP</td>
<td>Laparotomy</td>
</tr>
<tr>
<td>8</td>
<td>APPY</td>
<td>Appendix surgery</td>
</tr>
<tr>
<td>9</td>
<td>HER</td>
<td>Herniorrhaphy</td>
</tr>
<tr>
<td>10</td>
<td>NEPH</td>
<td>Kidney surgery</td>
</tr>
<tr>
<td>11</td>
<td>VHYS</td>
<td>Vaginal Hysterectomy</td>
</tr>
<tr>
<td>12</td>
<td>SPLE</td>
<td>Spleen surgery</td>
</tr>
<tr>
<td>13</td>
<td>CHOL</td>
<td>Gall bladder surgery</td>
</tr>
<tr>
<td>14</td>
<td>OVRY</td>
<td>Ovarian surgery</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Priority</th>
<th>Procedure Category</th>
<th>Neurosurgical (Brain/Spine) Operations</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>VSHN</td>
<td>Ventricular shunt</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>Procedure Category</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>
OPC-SSI Denominator for Procedure Reporting

Denominator Data

a) For each patient having at least one of the procedures included in the NHSN Operative Procedure category(s) for which SSI surveillance is being performed during the month, complete the Outpatient Procedure Component (OPC) Denominator for Procedure Form (CDC 57.404). The data are collected individually for each operative procedure category performed during the month specified on the OPC Monthly Reporting Plan. The Instructions for Completion of Outpatient Procedure Component (OPC) Denominator for Procedure Form (CDC 57.404) include brief instructions for collection and entry of each data element on the form.

b) Conduct post-discharge surveillance according to a formal active surveillance process. See Appendix A for the Post-discharge Surveillance Toolkit.

c) The surveillance period for a superficial SSI is 30 days after the procedure for all procedure categories. The surveillance period for deep and organ/space SSI is either 30 or 90 days, depending on the procedure category, as instructed in Table 2, Surveillance Periods for SSIs Following Selected NHSN Operative Procedure Categories.

d) Complete the Outpatient Procedure Component (OPC) Surgical Site Infection (SSI) Event Form (CDC 57.405) for each patient meeting the NHSN criteria for SSI, as defined in Surgical Site Infection Criteria, Table 4A for all procedures except breast & Table 4B for breast surgery procedures.

Table 7: Denominator Reporting Instructions

Denominator for procedure reporting instructions are guidelines for reporting data of each individual procedure that is to be counted (included) in the denominator of the selected procedure category. The instructions assist with maintaining data quality.

<table>
<thead>
<tr>
<th>Topic</th>
<th>Reporting Instruction</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>1. Wound class:</strong></td>
<td>A high wound class is not an exclusion for denominator reporting. If the procedure meets the definition of an NHSN operative procedure it should be reported in the denominator data regardless of wound class. NHSN will use the wound class for risk adjustment, as appropriate.</td>
</tr>
<tr>
<td><strong>2. Different operative procedure categories performed during same trip to the OR:</strong></td>
<td>If procedures in more than one NHSN operative procedure category are performed during the same trip to the operating room through the same or different incisions, an Outpatient Procedure Component (OPC) Denominator for Procedure Form (CDC 57.404) is reported for each procedure performed in the NHSN operative procedure category being monitored. For example: If a patient has an open reduction of fracture (FX) and knee arthroplasty (KPRO) performed during the same trip to the operating room and both procedure categories are being monitored and are included in the Monthly Reporting Plan,</td>
</tr>
<tr>
<td>Topic</td>
<td>Reporting Instruction</td>
</tr>
<tr>
<td>-------</td>
<td>-----------------------</td>
</tr>
<tr>
<td>complete an <a href="https://www.cdc.gov/nhsn/pdfs/ssi/OPC57404Denominator.pdf">Outpatient Procedure Component (OPC) Denominator for Procedure Form (CDC 57.404)</a> for each procedure.</td>
<td></td>
</tr>
</tbody>
</table>
| 3. **Duration** of the procedure when procedures from *more than one NHSN operative procedure category* is performed through the **same incision on the same trip to the OR**: | If more than one NHSN operative procedure category is performed through the same incision during the same trip to the operating room, record the combined duration of all procedures, which is the time from procedure/surgery start time to procedure/surgery finish time.  
For example:  
If a COLO and CHOL procedures are done through the same incision, the time from start time to finish time is reported for both operative procedures. |
| 4. **Duration** of operative procedures if patient has **two different NHSN operative procedures** performed via **separate incisions on the same trip to the OR**: | Try to determine the correct duration for each separate procedure (if this is documented), otherwise, take the time for both procedures and split it evenly between the two. |
| 5. **Same NHSN operative procedure category** via the **same incision/laparoscopic incision**, but different CPT codes during same trip to the OR: | If procedures of different CPT codes from the same NHSN operative procedure category are performed through the **same incision/laparoscopic sites**, record only one procedure for that category.  
For example:  
If a facility is performing surveillance for laminectomy procedures (LAM) and a patient undergoes a lumbar fusion of a couple contiguous vertebrae via one incision during the same trip to the operating room, complete one LAM [Outpatient Procedure Component (OPC) Denominator for Procedure Form (CDC 57.404)](https://www.cdc.gov/nhsn/pdfs/ssi/OPC57404Denominator.pdf) - because both procedures are in the LAM operative procedure category. |
| 6. **Same NHSN operative procedure category** via **separate incisions** during same trip to the OR: | For operative procedures that can be performed via **separate incisions** during same trip to operating room (specifically the following, AMP, BRST, CEA, FUSN, FX, HER, HPRO, KPRO, LAM, NEPH, OVRY, PVBY), separate [Outpatient Procedure Component (OPC) Denominator for Procedure Form (CDC 57.404)](https://www.cdc.gov/nhsn/pdfs/ssi/OPC57404Denominator.pdf) are completed. To document the duration of the procedures, indicate the procedure/surgery start time to procedure/surgery finish time |
### Reporting Instruction

<table>
<thead>
<tr>
<th>Topic</th>
<th>Reporting Instruction</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>for each procedure separately or, alternatively, take the total time for the procedures and split it evenly between procedures.</td>
</tr>
<tr>
<td><strong>NOTES:</strong></td>
<td></td>
</tr>
<tr>
<td>• A COLO procedure with a colostomy formation is entered as one COLO procedure.</td>
<td></td>
</tr>
<tr>
<td>• 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. Open (non-laparoscopic) hernia repairs are reported as one procedure for each hernia repaired via a separate incision, (specifically, 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.</td>
<td></td>
</tr>
</tbody>
</table>

7. **Patient expires in the OR:**

If a patient expires in the operating room, do not complete an Outpatient Procedure Component (OPC) Denominator for Procedure Form (CDC 57.404). This operative procedure is excluded from the denominator.

8. **HYST or VHYS:**

For the purpose of NHSN OPC-SSI reporting, hysterectomy procedures that involve an incision made into the abdomen, including trocar insertion, are included in the abdominal hysterectomy (HYST) category. The correct CPT hysterectomy procedure codes should be assigned by a medical record coder using current coding guidelines and conventions.

### Post-discharge Surveillance

When using OPC-SSI criteria for surveillance the method used for post-discharge SSI surveillance is a required element for reporting. NHSN require facilities to use a post-discharge surveillance process which is active and patient-based for identifying and detecting of SSIs events. An active surveillance process ensures that SSI events are associated with the correct NHSN operative procedure and are accurately attributed to the facility in which the procedure was performed. Post-discharge surveillance should include the full surveillance period for the given operative procedure category as listed in Table 2. See Appendix A for the NHSN OPC Post-discharge Surveillance Toolkit.

**Active post-discharge surveillance**
Active surveillance is a process in which the facility has a formal and routine process of identifying, investigating and detecting infections during the defined surveillance period. Active post-discharge surveillance may include but is not limited to:

- post-discharge letters or phone calls to patients
- inter-facility notification of patient encounters or admission
- review of medical or surgical clinic patient records
- post-discharge surgeon survey with listing of operative procedures performed

Any combination of these methods (or others identified by the facility) is acceptable for use to identify all SSIs; however, NHSN OPC-SSI criteria must be used. To minimize the workload of denominator data entry, upload of these data into the NHSN application is available using a comma-separated values (.csv) file. Instruction for .csv import can be found at https://www.cdc.gov/nhsn/pdfs/opc/importing-opc-procedure-data-508.pdf.

**Passive post-discharge surveillance**
Passive surveillance is a process which may include incidental or unsolicited post-discharge notifications of infections by surgeons, patients, family members or another facility. While passive surveillance may be an inherent part of post-discharge surveillance, it should not be the only process used for case identification.

If the facility already has an active standardized SSI surveillance process in place that is successfully identifying patients with infections post-discharge and is obtaining information from surgeons about potential SSIs, the facility may continue to use that process as long as the requirements of the OPC-SSI Protocol are met.

**Surveillance Reminders:**

- An SSI event is attributed to the facility in which the NHSN operative procedure was originally performed.
- Facilities that have identified potential SSI events that are attributable to procedures performed at a different facility should provide details of the potential events to the facility where the procedure was originally performed.

**Data Analyses**
Procedure (denominator) and SSI event (numerator) data that has been entered in to NHSN can be analyzed and visualized in a variety of reports.

**Types of SSI Analyses Reports:**

**Descriptive Analysis Reports**
Descriptive analysis options for numerator and denominator data such as line listings, frequency tables, and bar and pie charts are available in the NHSN application. These analysis options are also available to
analyze pathogens and antimicrobial susceptibility data reported for each SSI. NHSN Quick Reference. Guides on these reports can be found at https://www.cdc.gov/nhsn/ambulatory-surgery/ssi/index.html.

**SSI Standardized Infection Ratio (SIR) Reports**

The Standardized Infection Ratio (SIR) is calculated by dividing the number of observed infections by the number of predicted infections

\[
SIR = \frac{\text{Observed (O) HAI}s}{\text{Predicted (P) HAI}s}
\]

The SIR will be calculated only if the number of predicted HAIs (“numPred” in the NHSN application) is ≥ 1 to help enforce a minimum precision criterion.

The number of predicted infections is calculated using SSI probabilities estimated from multivariate logistic regression models constructed from NHSN data during a baseline time period, which represents a standard population’s SSI experience. Adult and pediatric procedures/SSIs are modeled separately; pediatric models will be available in the future. SSIs are included in the numerator of an SIR based on the date the procedure is preformed and not the date the event is identified. This is because the procedure carries the risk for the infection/SSI.

**Inclusion and Exclusion Criteria**

The OPC SSI SIR is calculated for facilities enrolled in NHSN as an Ambulatory Surgery Center (ASC). There is one SIR model available for outpatient adult procedures (and associated SSIs). Below is a summary of the OPC SSI SIR Model.

<table>
<thead>
<tr>
<th>OPC SSI SIR Model</th>
<th>Inclusion Criteria</th>
<th>Patient Population</th>
</tr>
</thead>
<tbody>
<tr>
<td>All SSI SIR Model</td>
<td>• Include only ambulatory surgery center procedures</td>
<td>Procedures in adult patients</td>
</tr>
<tr>
<td></td>
<td>• Include Superficial, Deep &amp; Organ/Space SSIs</td>
<td></td>
</tr>
<tr>
<td></td>
<td>• Superficial &amp; Deep Incisional SSIs are limited only</td>
<td></td>
</tr>
<tr>
<td></td>
<td>to primary incisional SSIs</td>
<td></td>
</tr>
<tr>
<td></td>
<td>• Include SSIs identified on active and passive</td>
<td></td>
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<tr>
<td></td>
<td>surveillance</td>
<td></td>
</tr>
</tbody>
</table>

In addition to the above inclusion criteria, there is also a list of exclusion criteria that applies to the OPC All SSI SIR model. The list of exclusion criteria applies to both procedures and the associated SSI events. Often the reason for excluding procedures and SSI events from the SIR calculation is due to potential data quality issues. It is important that facilities review their data for quality assurance and to determine the reason for exclusion from the SIR calculation.

More detailed information can be found in the NHSN Guide to the SIR: https://www.cdc.gov/nhsn/pdfs/ps-analysis-resources/nhsn-sir-guide.pdf.
NHSN Group Analysis
NHSN Group Users can perform the same analysis as facility level users in NHSN. Two helpful NHSN User Group report are the Line Listing - Membership Rights report and the Line Listing - Participation Alerts reports.

- The Line Listing - Membership Rights Report describes the rights conferred by each facility in the group. This report is helpful when determining the level and access to data in NHSN for a group.
- The Line Listing - Participation Alerts Report describes the unresolved NHSN alerts by alert type and facility. NHSN alerts serve as data reporting reminders and describe data quality issues. It is important to generate datasets in NHSN after alerts are resolved.

Resources for NHSN Group Users
This NHSN website contains guides that describe how to create a group, set up a confer rights template, and how to analyze group reports along with other group related topics: https://www.cdc.gov/nhsn/group-users/index.html.

Group User’s Guide to the “Line Listing- Participation Alerts” Report Option, this report is important as Groups may help their facilities with education about reporting data. The NHSN participation alerts help inform facilities about reporting data and data quality issues. The participation alerts line listing report is a tool that Groups can use to identify unresolved alerts for facilities that they have confer rights for in their group: https://www.cdc.gov/nhsn/pdfs/ps-analysis-resources/group-alerts.pdf.
References


Appendix A: Post-discharge Surveillance Toolkit

This toolkit was developed by NHSN to assist facilities in implementing an effective post-discharge surgical site infection surveillance process.

Contents:
The toolkit contains samples of a: Sample Letter, Post-discharge SSI Worksheet and Procedure Line List by Surgeon, along with instructions and helpful suggestions.

NOTE: If the facility already has an active standardized SSI surveillance process in place that is successfully identifying patients with infections post-discharge and is obtaining information from surgeons about potential SSIs, the facility may continue to use that process as long as the requirements of this Post-Discharge Surveillance Toolkit are met.

Instructions:
Based on the NHSN OPC-SSI Protocol, operative procedures must be followed for either a 30- or 90-day surveillance period after the operation in order to identify a potential SSI (Table 2).

1. Sample Letter – introduces the receiving surgeon and office staff to your facility’s post-discharge SSI surveillance program. It provides instructions and contact information if questions arise.

2. Procedure Line List by Surgeon - is line list that is generated at the end of every month (or 90-day period for select procedures). The line list will provide surgeons with a detailed list of each procedure they performed at the facility during the previous 30 (or 90) days.

3. SSI Worksheet – is used to allow surgeons or their designee to document whether any of their patients developed a suspected superficial, deep, or organ/space surgical site infection. This is a generic worksheet that can be used for any surgical procedure monitored by the facility.

The Procedure Line List and the Post-discharge SSI Worksheet can be sent to surgeons’ offices at the end of every surveillance period (30 or 90 days). Using the Procedure Line List as a guide, surgeons will complete one Worksheet for each patient who developed an SSI. All completed Worksheets should be sent back to the appropriate ASC staff to confirm that the documented SSI(s) correctly meets NHSN criteria. If the SSI(s) is confirmed, the infections must be entered into NHSN.

Instructions for the office staff on how to complete the Post-discharge SSI Worksheets can be customized based on your facility’s preferences.

IMPORTANT POINTS:

- Your facility must include either a Surgeon Code or Surgeon Name for each procedure entered in NHSN in order to generate the Procedure Line List by surgeon.
- The Procedure Line List and the SSI Worksheets should not be mailed until at least 30 or 90 days after the last surgical procedure so that the correct time period following the surgery has lapsed.
SAMPLE: LETTER
[Insert Name Ambulatory Surgery Center]     [Insert Date]
Post-discharge Surgical Site Infection Surveillance

Dear Office Staff,

Our records show that [Surgeon’s Name] performed surgical procedures at our facility during the [Insert Months & Year or surveillance period].

We are requesting your assistance with our post-discharge surgical site infection surveillance. Please review your records for each patient included on the line list.

- If a patient did not develop any surgical site infection check the “No Evidence of SSI box.”

- If a patient developed any signs or symptoms of infection, please complete the enclosed “Post-discharge Surgical Site Infection Worksheet.”

**NOTE:** Please make enough copies of the blank Post-discharge Surgical Site Infection Worksheet so that one worksheet can be completed for each patient with an SSI.

- Return this line list and any completed worksheets by [Insert Due Date]

The completed SSI worksheets and line list can be sent back via fax or mail. If you have any questions, please feel free to call.

Thank you for your assistance in ensuring our compliance with post-discharge SSI surveillance.

[Insert Name]
[Facility Name]
[Facility Address]
FAX: 000-000-0000
Phone: 000-000-0000
**SAMPLE: LINELIST for [Surgeon’s Name]**

[Insert Name Ambulatory Surgery Center]  
[Insert Date]

Post-discharge Surgical Site Infection Surveillance

<table>
<thead>
<tr>
<th>Patient Last Name</th>
<th>Patient First Name</th>
<th>Date of Birth</th>
<th>Gender</th>
<th>Procedure ID</th>
<th>Procedure Date</th>
<th>Procedure Category</th>
<th>Surgeon Code</th>
<th>No Evidence of SSI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Smith</td>
<td>Roger</td>
<td>10/20/1944</td>
<td>F</td>
<td>27467</td>
<td>06/30/2019</td>
<td>COLO</td>
<td>0103</td>
<td></td>
</tr>
<tr>
<td>Greene</td>
<td>Rachel</td>
<td>07/27/1949</td>
<td>F</td>
<td>27486</td>
<td>06/16/2019</td>
<td>COLO</td>
<td>0103</td>
<td></td>
</tr>
<tr>
<td>Blakeman</td>
<td>Mark</td>
<td>12/01/1927</td>
<td>M</td>
<td>27497</td>
<td>06/30/2019</td>
<td>COLO</td>
<td>0103</td>
<td></td>
</tr>
<tr>
<td>Fields</td>
<td>Rebecca</td>
<td>01/15/1960</td>
<td>F</td>
<td>27525</td>
<td>06/31/2019</td>
<td>COLO</td>
<td>0103</td>
<td></td>
</tr>
<tr>
<td>Hunter</td>
<td>Sean</td>
<td>09/23/1933</td>
<td>M</td>
<td>27531</td>
<td>06/24/2019</td>
<td>COLO</td>
<td>0103</td>
<td></td>
</tr>
<tr>
<td>Smith</td>
<td>Mary</td>
<td>07/16/1970</td>
<td>F</td>
<td>35014</td>
<td>06/09/2019</td>
<td>HYST</td>
<td>0103</td>
<td></td>
</tr>
<tr>
<td>Jones</td>
<td>SeQuisha</td>
<td>06/29/1972</td>
<td>F</td>
<td>35015</td>
<td>06/02/2019</td>
<td>HYST</td>
<td>0103</td>
<td></td>
</tr>
<tr>
<td>Archin</td>
<td>Latoya</td>
<td>09/03/1967</td>
<td>F</td>
<td>35016</td>
<td>06/07/2019</td>
<td>HYST</td>
<td>0103</td>
<td></td>
</tr>
</tbody>
</table>

To generate a line list from NHSN, see analysis resources found at [https://www.cdc.gov/nhsn/ambulatory-surgery/ssi/index.html](https://www.cdc.gov/nhsn/ambulatory-surgery/ssi/index.html).
# Post-discharge Worksheet for Suspected SSI

**[Insert Name Ambulatory Surgery Center]**

**[Insert Date]**

**Post-discharge Surgical Site Infection Surveillance**

## Patient Demographics:

<table>
<thead>
<tr>
<th>Patient Name (Last, First):</th>
</tr>
</thead>
<tbody>
<tr>
<td>Primary CPT Code of Procedure:</td>
</tr>
</tbody>
</table>

**Date SSI Identified:**

**Was the SSI identified on admission to a hospital?** Y N

*If Yes, name of facility: ________________________________

## Select the infection type and associated criteria (if known) from the options below:

- **□ A. Superficial Incisional SSI**: Involves only the skin and subcutaneous tissue of the incision

  **Criteria met (check all that apply):**
  - □ Purulent drainage from the superficial incision
  - □ Organisms identified from an aseptically obtained specimen from the superficial incision or subcutaneous tissue
  - □ Superficial incision that is deliberately opened by a surgeon, physician or physician designee and culture or non-culture based microbiologic testing is not performed.

  *If checked, please answer the following (check all that apply):*
  - ○ Pain or tenderness
  - ○ Localized swelling
  - ○ Redness (erythema)
  - ○ Heat
  - □ Diagnosis of a superficial incisional SSI by a physician or physician designee.

- **□ B. Deep Incisional SSI**: Involves deep soft tissues (for example, fascia and muscle layers)

  **Criteria met (check all that apply):**
  - □ Purulent drainage from the deep incision
  - □ Deep incision spontaneously dehisces, or is deliberately opened or aspirated by a surgeon, physician or physician designee and organism is identified from specimen or microbiologic testing not performed.

  *If checked, please answer the following (check all that apply):*
  - ○ Fever (>38°C)
  - ○ Localized pain or tenderness
  - □ Abscess or other evidence of infection involving the deep incision that is detected on gross anatomical or histopathologic exam, or imaging test
| C. Organ/Space: Involves any part of the body, (excluding skin incision, fascia, and muscle layers), that is opened or manipulated during the operative procedure |

**Criteria met (check all that apply):**
- Purulent drainage from a drain that is placed into the organ/space
- Organisms isolated from an aseptically-obtained specimen of fluid or tissue in the organ/space\(^1\)
- Abscess or other evidence of infection involving the organ/space that is detected on gross anatomical or histopathologic exam, or imaging test evidence consistent with infection

\(^1\) Culture or non-culture based microbiologic testing method.
\(^2\) Should be interpreted to mean surgeon(s), infectious disease physician, emergency physician, other physician on the case, or physician's designee (nurse practitioner or physician’s assistant).

**Additional comments:**

**Signature:**

**Date:**
Outpatient Procedure Component (OPC) Key Terms

**Ambulatory Surgery Center (ASC):** Based on the Code of Federal Regulations 42 CFR § 416.2, is any distinct entity that operates exclusively for the purpose of providing surgical services to patients not requiring hospitalization and in which the expected duration of services would not exceed 24 hours following an admission. The entity must have an agreement with CMS to participate in Medicare as an ASC and must meet the conditions set forth in 42 CFR § 416.

**Discharge:** A patient is considered discharged when they leave the confines of the outpatient facility.

**Encounter:** Any patient visit to an outpatient location where the patient completes the registration process upon entry into the facility. Some outpatient facilities also refer to this as admission into the facility.

**Hospital Outpatient Department (HOPD):** Based on the Code of Federal Regulations 42 CFR 440.20 is a location that provides outpatient hospital services, meaning preventive, diagnostic, therapeutic, rehabilitative, or palliative services that:

1. Are furnished to outpatients;
2. Are furnished by or under the direction of a physician or dentist; and
3. Are furnished by an institution that—
   i. Is licensed or formally approved as a hospital by an officially designated authority for State standard-setting; and
   ii. Meets the requirements for participation in Medicare as a hospital; and
4. May be limited by a Medicaid agency in the following manner: A Medicaid agency may exclude from the definition of “outpatient hospital services” those types of items and services that are not generally furnished by most hospitals in the State.

**Hospital Transfer or Admission:** Any transfer or admission from an ASC or HOPD directly to an acute care hospital including the hospital emergency room. Directly means upon discharge from the outpatient facility. This measure applies regardless of the reason for the hospital transfer/admission, and no direct hospital transfers/admissions should be excluded based on an assessment about whether the transfer/admission is or is not related to the outpatient facility admission. This measure excludes patients who are discharged from the outpatient facility and then later go to a hospital emergency room or acute care hospital, even if they do so on the same date as the outpatient facility admission.

**Operative procedure category:** A set of surgical procedures grouped together based on the NHSN definition of the procedure performed. Each Operative Procedure Category has been mapped with a list of CPT codes with procedure descriptions that aligns with the procedure category definition.

**Burn:** Any unintended tissue injury that occurs prior to discharge from the outpatient facility and is caused by any of the six recognized mechanisms: scalds, contact, fire, chemical, electrical or radiation.
Examples of devices that can cause burns include warming devices, prep solutions, electrosurgical units or lasers.

**Fall:** A sudden, uncontrolled, unintentional, and downward displacement of the body to the ground or other object, and that occurs within the confines of the outpatient facility. This definition excludes falls resulting from violent blows or other purposeful actions. It also excludes falls that do not occur within the confines of the outpatient facility, such as in a parking lot.

**Surgical site infection:** An infection meeting NHSN SSI criteria. The operative procedure must be one that is included in one of the NHSN operative procedure categories and occurs within a defined timeframe. There are three categories of SSI that are defined by CDC. Each SSI definition is related to the tissue depth of the infection, as illustrated below.

**Surveillance Period:** The timeframe following a NHSN operative procedures where you monitor and identify SSIs. The surveillance period is determined by the NHSN operative procedure category (for example, BRST-Breast surgery - 90-day surveillance period and HYST-abdominal hysterectomy – 30-day surveillance period). See Table 3

**Wrong** (site, side, patient, procedure, or implant): A procedure performed in such a way that was not in accordance with the intended procedure - specifically, wrong body site (part), wrong side of the body, wrong patient, wrong procedure and/or wrong implant. The procedure performed was not consistent with the informed consent signed by the patient/family/designee. May also be referred to as Wrong Event.
References
