

## BLOOD STREAM INFECTION (BSI)

(Revised January 1, 2016)

**Healthcare-associated infections (HAI):** All NHSN site specific infections must first meet the HAI definition as defined in the “Additional Information” checklist before a site specific infection (e.g., CLABSI) can be reported to NHSN.

**Present on Admission (POA):** Infections that are POA, as defined in the “Additional Information” checklist, are not considered HAIs and therefore are never reported to NHSN.

**Primary bloodstream infections (BSI):** Laboratory-confirmed bloodstream infections (LCBI) that are **not** secondary to an infection at another body site (see “Secondary Bloodstream Infection [BSI] Guide” at the end of this checklist as well as the checklist corresponding to the other body site).

**Date of event (DOE):** The BSI date of event is the date when the first element used to meet the laboratory-confirmed bloodstream infection (LCBI) criterion occurs for the first time within the 7- day infection window period. Synonym: infection date.

**Central line:** An intravascular catheter that terminates at or close to the heart or in one of the great vessels which is used for infusion, withdrawal of blood, or hemodynamic monitoring. The following are considered great vessels for the purpose of reporting central-line BSI and counting central-line days in the NHSN system:

Aorta	Brachiocephalic veins	Common iliac veins
Pulmonary artery	Internal jugular veins	Femoral veins
Superior vena cava	Subclavian veins	In neonates – the umbilical artery/vein
Inferior vena cava	External iliac veins	

### **NOTES:**

1. Neither the insertion site nor the type of device may be used to determine if a line qualifies as a central line. The device must terminate in one of the great vessels or in or near the heart, and be used for one of the purposes outlined above, to qualify as a central line.
2. At times an intravascular line may migrate from its original great vessel location. Subsequent to the original confirmation, NHSN does not require ongoing confirmation that a line resides in a great vessel. Therefore, once a line is identified to be a central line for NHSN purposes, it is considered a central line until discontinuation, regardless of migration, and associated central line days are included in any CLABSI surveillance being performed in that location.

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3. An introducer is considered an intravascular catheter, and depending on the location of its tip and use, may be a central line.
4. Pacemaker wires and other non-lumened devices inserted into central blood vessels or the heart are not considered central lines, because fluids are not infused, pushed, nor withdrawn through such devices.
5. The following devices are not considered central lines:
  - i. Extracorporeal membrane oxygenation (ECMO)
  - ii. Femoral arterial catheters
  - iii. Intra-aortic balloon pump (IABP) devices
  - iv. Hemodialysis reliable outflow (HeRO) dialysis catheters
  - v. Impella heart devices

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

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

**Temporary central line:** A non-tunneled, non-implanted catheter. **Permanent central line:** Includes

- Tunneled catheters, including certain dialysis catheters
- Implanted catheters (including ports)

### **Central line-associated BSI (CLABSI):**

○ where Patient had **ALL** **△** of the following:

- △** a laboratory-confirmed bloodstream infection (LCBI)
- △** central line (CL) or umbilical catheter (UC) was in place for >2 calendar days on the day of event (with day of device placement being Day 1)
- △** a CL or UC was in place on the date of event or the day before

**BLOOD STREAM INFECTION (BSI)****NOTE:**

- If a CL or UC was in place for >2 calendar days and then removed, the date of event of the LCBI must be the day of discontinuation or the next day to be a CLABSI.
- If the patient is admitted or transferred into a facility with an implanted central line (port) in place, and that is the patient's only central line, day of first access in an inpatient location is considered Day 1.
- "Access" is defined as line placement, infusion, or withdrawal through the line.
- Such lines continue to be eligible for CLABSI once they are accessed until they are either discontinued or the day after patient discharge (as per the Transfer Rule). Note that the "de-access" of a port does not result in the patient's removal from CLABSI surveillance.

**EXAMPLES of Determining a CLABSI vs. BSI:**

- Patient has central line inserted on June 1. On June 3, the central line is still in place and the patient's blood is collected for culture. The culture is positive for *S. aureus*. This is a CLABSI because the central line was in place for >2 calendar days (June 1, 2, and 3), and still in place, on the date of event (June 3).
- Patient has a central line inserted on June 1. On June 3, the central line is removed and on June 4 the patient's blood is collected for culture. The culture is positive with *S. aureus*. This is a CLABSI because the central line was in place for >2 calendar days (June 1, 2, and 3), and was in place the day before the date of event (June 4).
- Patient has a central line inserted on June 1. On June 3, the central line is removed. On June 5 patient spikes a fever of 38.3°C and the patient's blood is collected for culture. The culture is positive for *S. aureus*. This meets LCBI Criterion 1 but it is not a CLABSI because the Date of Event (June 5) did not occur on the day the central line was discontinued (June 3) nor the Next day (June 4).

**BLOOD STREAM INFECTION (BSI)****NOTE:**

- **Central lines that are removed and reinserted:** If, after central line removal, the patient is without a central line for at least one full calendar day (NOT to be read as 24 hours), then the central line day count to determine eligibility for a CLABSI, will start anew. If instead, a new central line is inserted before a full calendar day without a central line has passed, the central line day count, to determine eligibility for a CLABSI will continue uninterrupted. See figure below titled “Associating Central Lines (CL) Use to BSI”.
- Bloodstream infections will not be reported if they occur within the Repeat Infection Timeframe (RIT) of a previously identified BSI. See Repeat Infection Timeframe guidance in the “Additional Information” checklist.
- **Note that only primary BSIs create a BSI RIT. Secondary BSIs do not create a BSI RIT.**
- A positive blood specimen meeting LCBI criteria, that is accompanied by **documentation** of observed or suspected patient accession into vascular access lines, within the BSI infection window period, will be considered an LCBI, but not CLABSI for NHSN reporting purposes. A BSI RIT will be created. If reporting the BSI to NHSN, answer “No” to the risk factor event field “Central line?” If a facility is reporting CLABSI’s electronically to NHSN via Clinical Document Architecture (CDA), no CLABSI should be reported for this event, since this BSI is not considered associated to the central line. If blood specimens meeting LCBI criteria with a date of event outside the BSI RIT occur, they must be investigated as a part of any BSI surveillance. Documentation of observed or suspected patient accession into vascular access lines, within the BSI infection window period, will again be necessary in order to determine that the LCBI is not central-line associated for this reason.

**BLOOD STREAM INFECTION (BSI)****Associating Central Lines (CL) Use to BSI**

	March 31 (Hospital day 3)	April 1	April 2	April 3	April 4	April 5	April 6
Patient A	Central Line Day 3	Central Line Day 4	Central Line removed (CL Day 5)	Central Line replaced (CL Day 6)	Central Line Day 7	Central Line removed Day 8	No Central Line
Patient B	Central Line Day 3	Central Line Day 4	Central Line removed (CL Day 5)	No Central Line	Central Line replaced (CL Day 1)	CL Day 2	CL Day 3

**Rationale:**

NHSN surveillance for infection is not aimed at a specific device. Instead surveillance is aimed at identifying risk to the patient that is the result of device use in general.

- In the examples above, Patient A is eligible for a CLABSI beginning on March 31, through April 6, since a CL was in place for some portion of each calendar day until April 6. A BSI with date of event on April 6 would be a CLABSI since the CL had been in place > 2 days and was removed the day before the date of event.
- Patient B is eligible for a CLABSI on March 31 (CL Day 3) through April 3. The catheter had been in place > 2 days and an HAI occurring on the day of device discontinuation or the following calendar day is considered a device-associated infection. The patient is not eligible again for a CLABSI until April 6, when the second central line had been in place for > 2 days. (Note: NHSN will not require the BSI to be attributed to a specific central line when reporting.)

**Location of attribution:** The inpatient location where the patient was assigned on the date of the LCBI event, which is further defined as the date when the first element used to meet the LCBI criterion occurred (see Exception to Location of Attribution below).

**EXCEPTION TO LOCATION OF ATTRIBUTION:**

**Transfer Rule:** If the date of event for a CLABSI is the day of transfer or discharge, or the next day, the infection is attributed to the transferring location. Receiving facilities should share information about such HAIs with the transferring facility to enable

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reporting. This is called the Transfer Rule and examples are shown below:

- Patient with a central line in place in the SICU is transferred to the surgical ward. The day after transfer is the date of event for an LCBI. This is reported to NHSN as a CLABSI for the SICU.
- Patient with a central line in place is transferred from the medical ward to the coronary care ICU (CCU). An LCBI date of event is on day 4 in the CCU. The central line is still in place. This is reported to NHSN as a CLABSI for the CCU because the date of event was not the date of transfer from the medical ward, or the next day.
- After a two-week hospital stay, a patient on the urology ward of Hospital A has his only central line removed and is discharged home a few hours later. The IP from Hospital B calls the next day to report that this patient has been admitted to Hospital B and meets LCBI criteria. This CLABSI should be reported to NHSN for, and by, Hospital A and attributed to the urology ward because the date of event was the day of transfer.

	3/22	3/23	3/24
Locations in which patient was housed	Unit A	Unit A Unit B Unit C	Unit C Unit D This is also the date of event for a CLABSI. CLABSI is attributed to Unit A since Unit A was the first location in which the patient was housed the day before the date of event.

**BLOOD STREAM INFECTION (BSI)****INPATIENT DIALYSIS**

Inpatients receiving dialysis are included in any CLABSI surveillance in the location in which they are housed, regardless of whether or not the central line is the only central line and only accessed for dialysis. This also applies to patients in Long-Term Acute Care (LTAC) facilities within Acute Care Facilities when dialysis is received from the Acute Care Facility staff.

**EXAMPLES:** *CLABSIs in the following examples will be attributed to unit A*

- Patient on Unit A receives onsite dialysis by contracted dialysis staff
- Dialysis staff travels to Unit A to provide dialysis to Unit A patient
- Patient resides on Unit A for inpatient care, but is transported to dialysis unit within the facility for dialysis. Since CLABSIs cannot be attributed to non-bedded locations, such an event must be attributed to the inpatient location housing the patient.

**NOTE:**

Facilities may choose to capture information about the presence of a dialysis catheter in patients with LCBIs. The BSI collection form includes an optional data field “Any hemodialysis catheter present,” which may be marked yes or no, and used internally by facility to identify association of dialysis to LCBI

## BLOOD STREAM INFECTION (BSI)

### LCBI – Laboratory-Confirmed Blood Stream Infection

(Revised January 1, 2016)

**DEFINITION:** LCBI must meet at least **ONE** ☐ of the following criteria:

☐ **Criterion 1:** Patient has **ALL** ☐ of the following:

- ☐ Patient has a recognized pathogen identified from one or more blood specimens<sup>#</sup>

**AND**

- ☐ organism identified in blood is not related to an infection at another site. (See *Secondary Bloodstream Infection (BSI) Guide toward end of this checklist.*)

<sup>#</sup>by a culture or non-culture based microbiologic testing method which is performed for purposes of clinical diagnosis or treatment (e.g. not Active Surveillance Culture/Testing (ASC/AST)).

☐ **Criterion 2:** Patient has **ALL** ☐ of the following:

- ☐ Patient has at least **ONE** ☐ of the following signs or symptoms:

☐ fever (>38.0° C)

☐ chills

☐ hypotension

**AND**

- ☐ organism identified from blood is not related to an infection at another site. (See *Secondary Bloodstream Infection (BSI) Guide toward end of this checklist.*)

**AND**

- ☐ the same common commensal (example below) is identified from two or more blood specimens drawn on separate occasions <sup>#</sup> (See comment 5 )

*(Criterion elements must occur within the Infection Window Period (see Chapter 2), the 7-daytime period which includes the collection date of the positive blood culture, the 3 calendar days before and the 3 calendar days after.)*



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**EXAMPLES OF COMMON COMMENSALS\*:**

diphtheroids [ <i>Corynebacterium</i> spp., not <i>C. diphtheriae</i> ]	coagulase-negative staphylococci [including
<i>Propionibacterium</i> spp.	<i>Bacillus</i> spp. [not <i>B. anthracis</i> ]
viridans group streptococci	<i>Aerococcus</i> spp.
<i>Micrococcus</i> spp.	

\*See complete list of common commensals at <http://www.cdc.gov/nhsn/XLS/master-organism-Commensals-Lists.xlsx>

**NOTE:**

The matching common commensals represent a single element; therefore the collection date of the **first** common commensal is the date of the element used to determine the Date of the Event.

6/1/2014	6/2/2014	6/3/2014	6/4/2014	Date of LCBI Event = 6/1/2014
<i>S. epidermidis</i> (1 of 2)	<i>S. epidermidis</i> (2 of 2)	No LCBI elements	Fever >38	

<sup>#</sup>by a culture or non-culture based microbiologic testing method which is performed for purposes of clinical diagnosis or treatment (e.g. not Active Surveillance Culture/Testing (ASC/AST)).

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**Criterion 3:** Patient has **ALL** of the following:

○ Patient ≤1 year of age has at least **ONE** of the following signs or symptoms:

△ fever (>38°C)

△ hypothermia (<36°C)

△ apnea

△ bradycardia

**AND**

○ organism identified from blood is not related to an infection at another site.

(See *Secondary Bloodstream Infection (BSI) Guide* toward end of this checklist.)

**AND**

○ the same common commensal (example below) is identified from two or more blood specimens drawn on separate occasions<sup>#</sup> (See comment #5).

(Criterion elements must occur within the Infection Window Period (see Chapter 2), the 7-day time period which includes the collection date of the positive blood, the 3 calendar days before and the 3 calendar days after.)

### EXAMPLES OF COMMON COMMENSALS\*:

diphtheroids [ <i>Corynebacterium</i> spp., not <i>C. diphtheriae</i> ]	coagulase-negative staphylococci [including <i>S. epidermidis</i> ]
<i>Propionibacterium</i> spp.	<i>Bacillus</i> spp. [not <i>B. anthracis</i> ]
viridans group streptococci	<i>Aerococcus</i> spp.
<i>Micrococcus</i> spp.	

\* See complete list of common commensals at <http://www.cdc.gov/nhsn/XLS/master-organism-Commensals-Lists.xlsx>

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### **NOTE:**

The matching common commensals represent a single element; therefore the collection date of the **first** common commensal is the date of the element used to determine the Date of the Event.

6/1/2014	6/2/2014	6/3/2014	6/4/2014	Date of LCBI Event = 6/1/2014
S. epidermidis (1 of 2)	S. epidermidis (2 of 2)	No LCBI elements	Apnea documented	

# by a culture or non-culture based microbiologic testing method which is performed for purposes of clinical diagnosis or treatment (e.g. not Active Surveillance Culture/Testing (ASC/AST))

## BLOOD STREAM INFECTION (BSI)

**MBI-LCBI – Mucosal Barrier Injury Laboratory-Confirmed Blood Stream Infection**  
*(Revised January 1, 2016)*

**NOTE:** For MBI-LCBIs, ANC/WBC levels should not be used to set the IWP or to identify the date of event. MBI-LCBIs are subsets of LCBIs and therefore the date of the LCBI would be the date of the MBI-LCBI event.

**DEFINITION:** MBI-LCBI must meet at least **ONE** ☐ of the following criteria:

☐ **Criterion 1:** Patient has **ALL** ☐ of the following:

☐ Patient of any age has **BOTH**  of the following:

meets criterion 1 for LCBI with at least one blood specimen identified<sup>#</sup>

has at least one blood culture growing any of the following intestinal organisms with no other organisms isolated: (See Comment #6)

<i>Bacteroides</i> spp.	<i>Enterococcus</i> spp.	<i>Prevotella</i> spp.
<i>Candida</i> spp.	<i>Fusobacterium</i> spp.	<i>Veillonella</i> spp.
<i>Clostridium</i> spp.	<i>Peptostreptococcus</i> spp.	Enterobacteriaceae*

\*See table toward the end of this checklist for partial list of eligible *Enterobacteriaceae* genera.

**AND**

☐ Patient meets at least **ONE**  of the following:

is an allogeneic hematopoietic stem cell transplant recipient within the past year with **ONE** ☐ of the following documented during same hospitalization as positive blood specimen:

☐ grade III or IV gastrointestinal graft versus host disease (GI GVHD)

☐ ≥1 liter diarrhea in a 24-hour period (or ≥20 mL/kg in a 24-hour period for patients <18 years of age) with onset on or within 7 calendar days before the date the positive blood culture was collected

is neutropenic, defined as at least 2 separate days with **ONE** ☐ of the following:

☐ values of absolute neutrophil count (ANC) <500 cells/mm<sup>3</sup> within a 7-day time period which includes the date the positive blood specimen was collected (Day 1), the 3 calendar days before **AND** the 3 calendar days after. (See table titled Examples Illustrating the MBI-LCBI Criterion for Neutropenia)

☐ total white blood cell count (WBC) <500 cells/mm<sup>3</sup> within a seven-day time period which includes the date the positive culture was collected (Day 1), the 3 calendar days before **AND** the 3 calendar days after. (See table titled Examples Illustrating the MBI-LCBI Criterion for Neutropenia)

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- Criterion 2:** Patient has **ALL** of the following: Patient of any age has **BOTH** of the following:
- △ meets criterion 2 for LCBI with at least one blood specimen identified<sup>#</sup>
  - △ has at least one blood specimen that is growing only *viridans* group *streptococci* with no other organisms
- AND**
- Patient meets at least **ONE** of the following:
    - △ is an allogeneic hematopoietic stem cell transplant recipient within the past year with **ONE** of the following documented during same hospitalization as positive blood specimen:
      - grade III or IV gastrointestinal graft versus host disease (GI GVHD)
      - ≥1 liter diarrhea in a 24-hour period (or ≥20 mL/kg in a 24-hour period for patients <18 years of age) with onset on or within 7 calendar days before the date the first positive blood specimen was collected.
    - △ is neutropenic, defined as at least 2 separate days with **ONE** of the following:
      - values of absolute neutrophil count (ANC) <500 cells/mm<sup>3</sup> within a 7-day time period which includes the date the positive specimen was collected (Day 1), the 3 calendar days before **AND** the 3 calendar days after. (See table titled Examples Illustrating the MBI-LCBI Criterion for Neutropenia)
      - total white blood cell count (WBC) <500 cells/mm<sup>3</sup> within a 7-day time period which includes the date the positive specimen was collected (Day 1), the 3 calendar days before **AND** the 3 calendar days after. (See table titled Examples Illustrating the MBI-LCBI Criterion for Neutropenia)

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❑ **Criterion 3:** Patient has **ALL** ○ of the following:

○ Patient ≤1 year of age has **BOTH** △ of the following:

△ meets criterion 3 for LCBI with at least one blood specimen are identified<sup>#</sup>

△ has at least one blood specimen that is growing only *viridans group streptococci* with no other organisms isolated.

**AND**

○ Patient meets at least **ONE** △ of the following:

△ is an allogeneic hematopoietic stem cell transplant recipient within the past year with **ONE** □ of the following documented during same hospitalization as positive blood specimen:

□ grade III or IV gastrointestinal graft versus host disease (GI GVHD)

□ ≥20 mL/kg diarrhea in a 24-hour period with onset on or within 7 calendar days before the date the first positive blood specimen is collected.

△ is neutropenic, defined as at least 2 separate days with **ONE** □ of the following:

□ values of absolute neutrophil count (ANC) <500 cells/mm<sup>3</sup> within a 7-day time period which includes the date the positive culture was collected (Day 1), the 3 calendar days before **AND** the 3 calendar days after.

(See table titled Examples Illustrating the MBI-LCBI Criterion for Neutropenia)

□ total white blood cell count (WBC) <500 cells/mm<sup>3</sup> within a 7-day time period which includes the date the positive culture was collected (Day 1), the 3 calendar days before **AND** the 3 calendar days after.

(See table titled Examples Illustrating the MBI-LCBI Criterion for Neutropenia)

<sup>#</sup>by a culture or non-culture based microbiologic testing method which is performed for purposes of clinical diagnosis or treatment (e.g. not Active Surveillance Culture/Testing (ASC/AST)).

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### **Comments:** (Revised January 1, 2016)

1. A positive blood specimen meeting LCBI criteria, that is accompanied by **documentation** of observed or suspected patient accession into vascular access lines, within the BSI infection window period, will be considered an LCBI, but not CLABSI for NHSN reporting purposes. A BSI RIT will be created. If reporting the BSI to NHSN, answer “No” to the event field “central line?” If a facility is reporting CLABSIs electronically to NHSN via Clinical Document Architecture (CDA) no CLABSI should be reported for this event, since this BSI is not considered associated to the central line. If blood cultures collected after the BSI RIT are again positive, they must be investigated as part of any BSI surveillance. Documentation of observed or suspected patient accession into vascular access lines, within the BSI infection window period, will again be necessary in order to determine that the LCBI is not central-line associated.
2. In LCBI criterion 1, the term “recognized pathogen” includes any organism not included on the common commensal list (*See criteria 2 and 3 or Supporting Material section at <http://www.cdc.gov/nhsn/XLS/master-organism-Com-Commensals-Lists.xlsx> for the list of common commensals*).

#### Exceptions:

- *Salmonella spp.* are excluded as pathogens for LCBI. These organisms may be secondary BSIs but will not be reported as the sole pathogen in a primary BSI.
  - Organisms belonging to the following genera cannot be used to meet any NHSN definitions: *Blastomyces*, *Histoplasma*, *Coccidioides*, *Paracoccidioides*, *Cryptococcus* and *Pneumocystis*. These organisms are typically causes of community-associated infections and are rarely known to cause healthcare-associated infections and therefore are excluded.
3. LCBI criteria 1 and 2 and MCI-LCBI criteria 1 and 2 may be used for patients of any age, including those patients  $\leq 1$  year of age.
  4. In LCBI criteria 2 and 3, if the pathogen or common commensal is identified to the species level from one blood specimen, and a companion blood specimen is identified with only a descriptive name, which is complementary to the companion culture (e.g., to the genus level), then it is assumed that the organisms are the same. The organism identified to the species level should be reported as the infecting organism along with its antibiogram if available (**See table below titled “Examples of How to Report Speciated and Unspeciated Organisms Isolated from Blood Cultures.”**). Only genus and species identification should be used to determine the sameness of organisms (i.e., matching organisms). No additional comparative methods should be used (e.g., morphology or antibiograms) because

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laboratory testing capabilities and protocols may vary between facilities. This will reduce reporting variability, solely due to laboratory practice, between facilities reporting LCBI meeting criterion 2. Report the organism to the genus/species level only once, and if antibiogram data are available, report the results from the most resistant panel.

5. In LCBI criteria 2 and 3, the phrase “two or more blood specimens drawn on separate occasions” means, 1) that blood from at least two separate blood draws were collected on the same or consecutive calendar days, and 2) were collected in a manner which suggests that 2 separate blood draw site preparations were performed. This will reduce misidentification of contaminated blood specimens as LCBI. For example, blood specimens drawn from different sites (e.g., different venipunctures, a combination of venipuncture and lumen withdrawal, or different lumens of the same central line) should undergo separate decontaminations and are therefore considered drawn on “separate occasions”.
6. For pediatric patients, due to volume constraints, a blood specimen may consist of a single bottle. Therefore, to meet this part of the criterion, each bottle from two, single bottle blood draws would have to be positive for the same common commensal.
7. Specimen Collection Considerations: Although blood specimens drawn through central lines can have a higher rate of contamination than blood specimens collected through peripheral venipuncture<sup>3,4</sup>, all positive blood specimens, regardless of the sites from which they were collected, must be included when conducting in-plan CLABSI surveillance.
8. In MBI-LCBI 1, 2 and 3, “No other organisms” means there is no identification of a non-MBI-LCBI pathogens (e.g., *S. aureus*) or 2 matching common commensals (e.g., coagulase-negative staphylococci) collected from blood on separate occasions that would otherwise meet LCBI criteria. If this occurs, the infection should not be classified as MBI-LCBI.

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**REPORTING INSTRUCTIONS:** (Revised January 1, 2016)

1. Report organisms identified from blood as BSI–LCBI when no other site of infection is evident. Note: VASC infections with positive blood specimens should be reported as BSI-LCBI (see Reporting Instruction 5 below for exception)  
(see “*Secondary Bloodstream Infection (BSI) Guide*” found after the tables below.)
2. When another blood specimen is collected during the RIT of an identified MBI-LCBI, which is positive for an organism excluded from MBI-LCBI criteria, the MBI-LCBI event is edited to become an LCBI and the organism is added.
3. Catheter tip cultures are not used to determine whether a patient has a primary BSI.



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4. Purulent phlebitis confirmed with a positive semi quantitative culture of a catheter tip, **but with either negative or no blood culture** is considered a CVS-VASC, not an LCBI, SST-SKIN, or an SST-ST infection.
5. Occasionally, a patient with both a central IV line and another vascular access device develops a primary bloodstream infection (LCBI) that can clearly be attributed to the other vascular access site. If both pus at the insertion site and a culture of that pus, collected during the LCBI infection window period, has at least one matching organism to the blood specimen, the LCBI will not be considered associated with the central line. In this situation, enter "No" for the filed "Central line" in the NHSN application. You should, however, include the patient's central line days in the summary denominator count. Vascular access devices included in this exception are limited to:
  - Peripheral IV
  - Arteriovenous fistula
  - Arteriovenous graft
  - on-accessed central line (not accessed nor inserted during the hospitalization)

**Examples of How to Report Speciated and Unspeciated Organisms Identified from Blood Specimens**

Culture Report	Companion Culture Report	Report as...
<i>Coagulase-positive staphylococci</i>	<i>S. aureus</i>	<i>S. aureus</i>
<i>S. epidermidis</i>	Coagulase-negative staphylococci	<i>S. epidermidis</i>
<i>Enterococcus</i> spp.	<i>E. faecium</i>	<i>E. faecium</i>
<i>Bacillus</i> spp. (not <i>anthracis</i> )	<i>B. cereus</i>	<i>B. cereus</i>
<i>S. salivarius</i>	Strep viridans	<i>S. salivarius</i>

**Partial List of Criterion 1 MBI-LCBI Eligible Enterobacteriaceae Genera\***

<i>Citrobacter</i>	<i>Enterobacter</i>	<i>Escherichia</i>		<i>Klebsiella</i>
<i>Proteus</i>	<i>Providencia</i>	<i>Salmonella</i>		<i>Serratia</i>
<i>Shigella</i>	<i>Yersina</i>			

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\*See complete list of MBI pathogens by selecting the MBI Organisms tab at the bottom of the Excel worksheet at common commensals at <http://www.cdc.gov/nhsn/XLS/master-organism-Com-Commensals-Lists.xlsx>

### Examples Illustrating the MBI-LCBI Criteria for Neutropenia

		Day -7	Day -6	Day -5	Day -4	Day -3	Day -2	Day -1	Day 1*	Day 2	Day 3	Day 4
Pt. A	WBC	100	800	400	300	ND	ND	320	400 + BC* w/ <i>Candida</i> spp. x1	ND	550	600
Pt. B	ANC	ND	410	130	ND	ND	120	110	ND +BC* w/ <i>viridans</i> strep x2 and fever >38°C	110	300	320
Pt. C	WBC	100	800	400	300	ND	ND	ND	600 + BC* w/ <i>Candida</i> spp. x1	230	ND	400

ND = not done; \* Day the blood specimen that was positive was collected

Patient A meets MBI-LCBI criterion 1, sub-criterion 2: Positive blood specimen with intestinal organism (*Candida* spp.) and neutropenia (2 separate days of WBC <500 cells/mm<sup>3</sup> occurring on the date the positive blood specimen was collected [Day 1] or during the 3 days before or the 3 days after that date). In this case, the Day 1 value = 400, and Day -1 value = 320.

Patient B meets MBI-LCBI criterion 2, sub-criterion 2: At least two positive blood specimens with *viridans* group streptococci (in this case, two positive), and fever >38°C and neutropenia (two separate days of ANC <500 cells/mm<sup>3</sup> occurring on the date the positive blood culture was collected [Day 1] or during the 3 days before or the 3 days after that date). In this case, the Day -1 value = 110 and Day -2 value = 120.

**NOTE:** Any two of Days -2,-1, 2, 3, and 4 could be used to meet this requirement since WBC or ANC under 500 were present on those days.

Patient C meets MBI-LCBI criterion 1, sub-criterion 2: Positive blood specimen with intestinal organism (*Candida* spp.) and neutropenia (2 separate days of WBC <500 cells/mm<sup>3</sup> occurring on the date the positive blood specimen was collected [Day 1] or during the 3 days before or the 3 days after that date). In this case, Day 2 value =230 and Day 4 value = 400).

## BLOOD STREAM INFECTION (BSI)

### Secondary Bloodstream Infection (BSI) Guide (not applicable to Ventilator-associated Events [VAE])

(Revised January 1, 2016)

The purpose of using the CDC/NHSN infection criteria is to identify and consistently categorize infections that are healthcare-associated into major and specific infection sites or types. LCBI criteria include the caveat that the organism(s) identified from the blood cannot be related to infection at another site (i.e., must be a primary BSI). One must be sure that there is no other CDC-defined primary site of infection that may have seeded the bloodstream secondarily; otherwise the bloodstream infection may be misclassified as a primary BSI and erroneously associated with the use of a central line, i.e., called a CLABSI. For locations performing in-plan VAE surveillance refer to the [VAE chapter](#) for specific guidance on assigning a secondary BSI to a VAE.

#### Secondary BSI Scenarios

**For purposes of NHSN, in order for a bloodstream infection to be determined secondary to another site of infection the following requirements must be met:†**

- △ Must meet one △ of the NHSN site-specific definitions (CDC/NHSN Surveillance Definitions for Specific Types of Infections, UTI, PNEU or SSI)

**AND**

**ONE □** of the following:

1. □ Patient has both ○ of the following:
  - An organism identified from the site specific infection is used as an element to meet the site-specific infection criterion.
  - Blood specimen contains at least one matching organism to that site-specific specimen, and is collected during the secondary BSI attribution period
- OR**
2. □ The positive blood specimen is an element used to meet the site-specific infection criterion, and is collected during the site-specific infection's infection window period.

#### †Exception:

Necrotizing enterocolitis (NEC) criteria include neither a site specific specimen nor an organism identified from blood specimen, however an exception for assigning a BSI secondary to NEC is provided.

A BSI is considered secondary to NEC if the patient meets one of the two NEC criteria **AND** an organism identified from blood specimen collected during the secondary BSI attribution period is an LCBI pathogen or the same common commensal is identified from two or more blood specimens drawn on separate occasions on the same or consecutive days.

## BLOOD STREAM INFECTION (BSI)

### Secondary BSI Scenarios

Below are examples with guidance on how to distinguish between the primary or secondary nature of a BSI. The definition of “matching organisms”, and important notes and reporting instructions are also provided.

See the flow diagram at the end of the checklist titled: “Secondary BSI Guide,” for algorithmic display of the following instructions.

#### Scenario 1:

**An organism identified from the site-specific infection is used as an element to meet the site-specific infection criterion, AND the blood specimen contains at least one matching organism to that site specific specimen. The positive blood specimen must be collected during the site-specific infection’s secondary BSI attribution period.**

#### Examples:

- a. Patient meets NHSN criteria for a symptomatic urinary tract infection (suprapubic tenderness and  $>10^5$  CFU/ml of *E. coli*) and blood specimen collected during the SUTI secondary BSI attribution period is positive for *E. coli*. This is a SUTI with a secondary BSI and the reported organism is *E. coli*.
- b. Patient meets NHSN criteria for a symptomatic urinary tract infection (suprapubic tenderness and  $>10^5$  CFU/ml of *E. coli*) and blood specimen collected during the SUTI secondary BSI attribution period grows *E. coli* and *P. aeruginosa*. This is a SUTI with a secondary BSI and the reported organisms are *E. coli* and *P. aeruginosa*, since both site and blood specimens are positive for at least one matching pathogen.
- c. Patient meets NHSN criteria for a symptomatic urinary tract infection (suprapubic tenderness and  $>10^5$  CFU/ml of *E. coli*) and a single blood specimen collected during the SUTI secondary BSI attribution period grows *E. coli* and *S. epidermidis*. This is a SUTI with a secondary BSI and the reported organism is only *E. coli*, since the single common commensal *S. epidermidis* positive blood specimen by itself does not meet BSI criteria.

## BLOOD STREAM INFECTION (BSI)

### **Scenario 2:**

The positive blood culture is an element used to meet the site-specific infection criterion, and is collected during the site specific infection's infection window period. (For your convenience, a list of infection criteria that include positive blood culture as an element are included in the table, "Site-specific criteria that require blood cultures").

### **Examples:**

- a. Patient becomes febrile and complains of nausea and abdominal pain. CT Scan done that day shows fluid collection suggestive of infection. Blood specimen collected that day results in identification of *Bacteroides fragilis*. Because the patient meets IAB criterion 3b, using the identification of an organisms from the blood specimen as an element (fever, nausea or abdominal pain, positive blood specimen and CT scan showing infection in abdominal cavity), the BSI is considered secondary to IAB.
- b. Patient is febrile, has a new onset of cough and has positive chest imaging test indicating the presence of an infiltrate. Blood specimens collected identify *Pseudomonas aeruginosa*. Because the patient can meet PNU 2 definition by using identification of organisms from blood specimen as one of the elements of the infection criterion (i.e., infiltrate on chest imaging test, fever, new onset of cough and organism identified from blood specimen), the BSI is considered secondary to PNEU.

**Note:** In scenarios where an NHSN infection definition can be met using more than one criterion of the infection definition, it is possible that identification of organism from blood and site-specific specimens may not match and a BSI may still be considered as a secondary BSI. Consider the following:

### **Examples:**

- During the SSI surveillance period, a postoperative patient becomes febrile and complains of nausea and abdominal pain. CT scan done that day shows fluid collection suggestive of infection. Culture results show *Escherichia coli* from the T-tube drainage specimen and the blood specimen grows *Bacteroides fragilis*. Although the organisms in the blood culture and site-specific culture do not match for at least one organism, the blood culture is considered secondary to IAB. This is because the patient meets organ/space SSI IAB criterion 3b, using the identification of organism in blood specimen as an element (fever, nausea or abdominal pain, organisms identified from blood specimen and CT scan showing infection in abdominal cavity). This patient also meets IAB criterion 3a using the positive site culture plus fever, and nausea or abdominal pain even though the organism involved is different from that used for IAB criterion 3b. In this case the

## BLOOD STREAM INFECTION (BSI)

BSI is considered secondary to the organ/space SSI IAB and both organisms would be listed as IAB infection pathogens.

- Patient is febrile, has a new onset of cough and has positive chest imaging test indicating the presence of an infiltrate. Blood and bronchoalveolar lavage (BAL) specimens are collected. Results identify *Klebsiella pneumonia*  $>10^4$  CFU/ml from the BAL and *Pseudomonas aeruginosa* from the blood. Although the organisms in the blood specimen and site-specific specimen do not match for at least one organism, because the patient can meet PNU2 definition using either the identification of organism from blood specimen or BAL specimen as one of the elements of the infection criterion (i.e. infiltrate on chest imaging test, fever, new onset of cough and organism identified from blood specimen or identified from BAL specimen), the blood is considered a secondary BSI to PNEU and both organisms would be listed as PNEU pathogens.

**If there is no matching organism identified from blood and site-specific specimen which is used to meet the site-specific infection definition, nor is an organism identified from blood specimen used to meet the site-specific infection criterion, secondary BSI attribution cannot be assigned.**

### Examples:

- a. Patient has pustules on their abdomen with tenderness and swelling. Purulent material is obtained from the pustules and is positive for *Streptococcus* Group B. A blood specimen collected the same day identifies methicillin resistant *Staphylococcus aureus*. Because the organisms from the site and blood specimens do not match, and there is no site-specific criterion for SKIN that includes organisms identified from blood specimen, both a site-specific infection, SKIN (criteria 1 and 2a) and a primary BSI would be reported.
- b. A patient has an abscess in the soft tissue around a percutaneous endoscopic gastrostomy (PEG) tube, identified by CT scan, and there is also purulent drainage from that site. No site-specific specimen was collected, but a blood specimen is positive for *Staphylococcus aureus*. No other sites of infection are identified. Because no culture of the site was collected, and the patient therefore cannot meet ST criterion 1, and because there is no ST criterion which uses identification of organism from blood specimen as an element, this patient has a ST infection with unknown pathogen (criterion 2), and a primary BSI with the pathogen *Staphylococcus aureus* for NHSN purposes.

## BLOOD STREAM INFECTION (BSI)

Site-specific criteria that require blood cultures

Organisms identified from blood as an element		
Site	Element	Page
Burn	1	17-23
IAB	2b	17-19
JNT	3c	17-6
MEN	2c & 3c	17-8
OREP	3a	17-22
PNU2	Lab finding	6-6
PNU3	Lab finding	6-8
UMB	1b	17-25

Organisms identified from blood <u>with</u> imaging test evidence of infection		
Site	Element	Page
Bone	3a	17-5
Disc	3a	17-5
GIT	2c	17-18
IAB	3b	17-19
SA	3a	17-9
USI	3b & 4b	17-26
ENDO	4a, 4b, 5a & 5b (specific organisms) 6e & 7e plus other criteria as listed	17-10



**BLOOD STREAM INFECTION (BSI)**

A **matching organism** is defined as one of the following:

1. If genus and species are identified in both specimens, they must be the same.

**Examples:**

- a. A blood specimen reported as *Enterobacter cloacae* and an intraabdominal specimen of *Enterobacter cloacae* are matching organisms.
  - b. A blood specimen reported as *Enterobacter cloacae* and an intraabdominal specimen of *Enterobacter aerogenes* are NOT matching organisms as the species are different.
2. If the organism is less definitively identified in one specimen than the other, the identifications must be complimentary.

**Examples:**

- a. A surgical wound growing *Pseudomonas* spp. and a blood specimen growing *Pseudomonas aeruginosa* are considered a match at the genus level and therefore the BSI is reported as secondary to the SSI.
- b. A blood specimen reported as *Candida albicans* and a culture from a decubitus reported as, "yeast not otherwise specified," are considered to have matching organisms because the organisms are complementary, i.e. *Candida* is a type of yeast.

**NOTES:** (Revised January 1, 2016)

1. Antibiograms of the blood and potential primary site isolates do not have to match.
2. If the blood specimen by itself does not meet BSI criteria (e.g., only one positive blood specimen positive for a common commensal), then that specimen may not be used to indicate the presence of a secondary BSI (see scenario 1c).

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**REPORTING INSTRUCTIONS:** (Revised January 1, 2016)

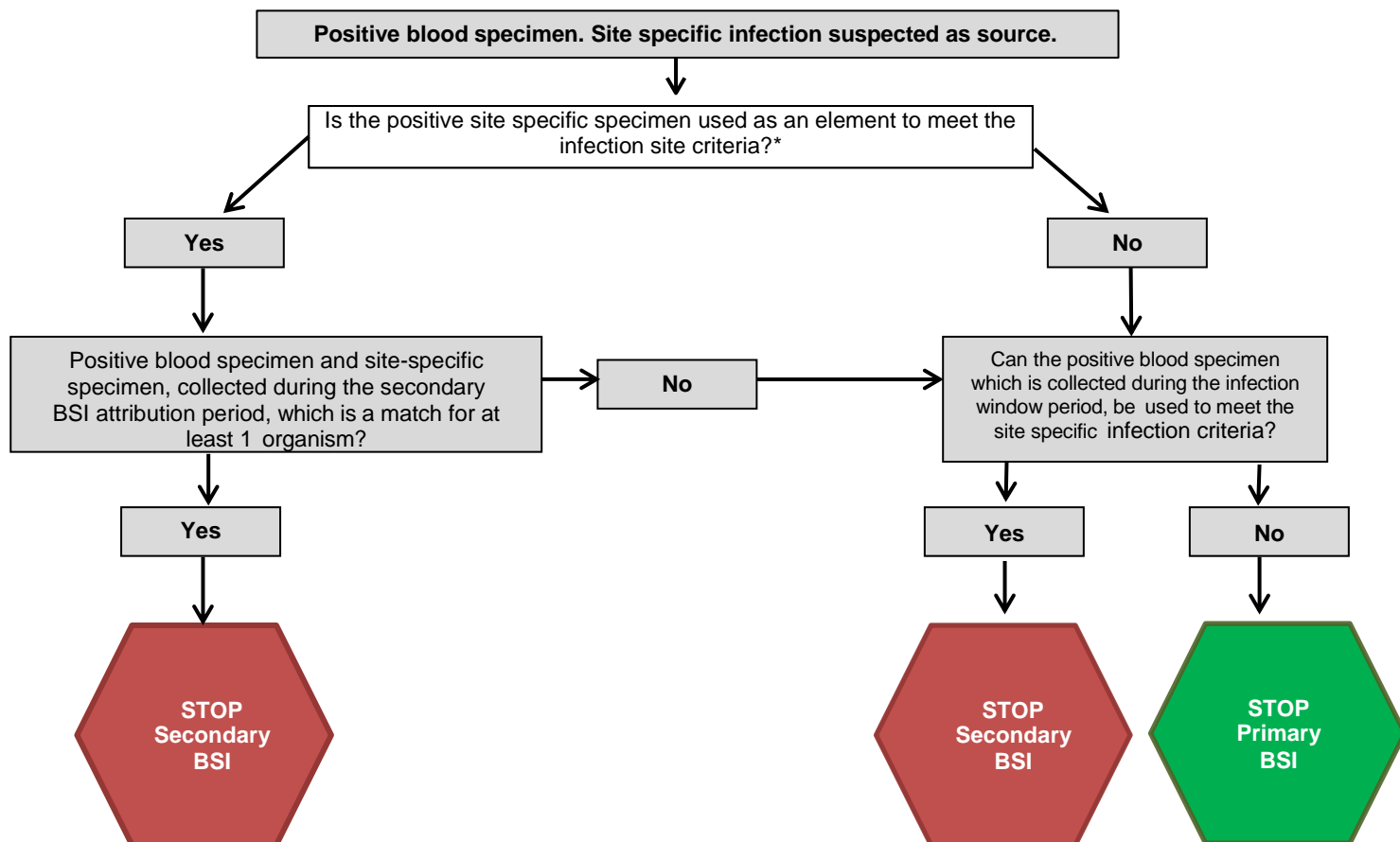
1. For reporting secondary BSI for possible VAP( PVAP), see [Chapter 10](#).
2. Do not report secondary bloodstream infection for vascular (VASC) infections, Ventilator-Associated Conditions (VAC), or Infection-related Ventilator-Associated Complications (IVAC), pneumonia 1 (PNEU 1).
3. When a BSI is suspected to be secondary to a lower respiratory tract infection and the BSI cannot be determined to be secondary to VAE, the PNEU definitions are available for secondary BSI assignment.
4. Site-specific organism exclusions apply to secondary BSI attribution as well.



**BLOOD STREAM INFECTION (BSI)**

**Secondary BSI Guide for Eligible Organisms\* \*\***  
**(Not applicable to Ventilator – Associated Events [VAE])**

(Revised January 2016)



**\*\*Exception:** Necrotizing enterocolitis (NEC) criteria include neither a site specific specimen nor organism identified from blood specimen, however an exception for assigning a BSI secondary to NEC is provided. A BSI is considered secondary to NEC if the patient meets one of the 2 NEC criteria AND an organism identified from blood specimen collected during the secondary BSI attribution period is an LCBI pathogen or the same common commensal is identified from 2 or more blood specimens drawn on separate occasions collected on the same or consecutive days.