

BLOODSTREAM INFECTIONS (BSI)

EPIDEMIOLOGY, PATHOPHYSIOLOGY, AND PREVENTION

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
Disclosures

Up-to-Date Royalties (Pelvic Osteomyelitis)
Legal Consultant (PJI and septic arthritis)

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Objectives



- Understand the impact of bloodstream infections
- Understand the incidence and causative pathogens of bloodstream infections
- Understand the risk factors for healthcare-associated bloodstream infections
- Understand how we define bloodstream infection, both clinically and epidemiologically
- Understand the prevention and control of bloodstream infections

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BSI: Impact on Healthcare

Approximately 72,000 primary bloodstream infections per year

- Accounts for ~10% of healthcare-associated infections (rank = 5)

Rate of BSIs varies by:

- Hospital size, unit, and service
- Population served (elderly/infants, acute/chronic)
- Use and type of intravascular access device
- Time-trends
- Endemic/Epidemic

Major risk = central venous catheter

- CLABSI associated with increased length of stay and increased cost (\$3,700 to \$39,000 per episode)

Magill SS, et al. New Engl J Med 2014;370:1198; Marschall J, et al. ICHE 2014;35:753-771

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Table 2. Distribution of 504 Health Care-Associated Infections.*

Type of Infection	Rank	No. of Infections	Percentage of All Health Care-Associated Infections (95% CI)
Pneumonia†	1 (tie)	110	21.8 (18.4–25.6)
Surgical-site infection	1 (tie)	110	21.8 (18.4–25.6)
Gastrointestinal infection	3	86	17.1 (14.0–20.5)
Urinary tract infection‡	4	65	12.9 (10.2–16.0)
Primary bloodstream infection§	5	50	9.9 (7.5–12.8)
Eye, ear, nose, throat, or mouth infection	6	28	5.6 (3.8–7.8)
Lower respiratory tract infection	7	20	4.0 (2.5–6.0)
Skin and soft-tissue infection	8	16	3.2 (1.9–5.0)
Cardiovascular system infection	9	6	1.2 (0.5–2.5)
Bone and joint infection	10	5	1.0 (0.4–2.2)
Central nervous system infection	11	4	0.8 (0.3–1.9)
Reproductive tract infection	12	3	0.6 (0.2–1.6)
Systemic infection	13	1	0.2 (0.01–1.0)

50 Primary BSI
42 (82%) CLABSI
37 Secondary BSI

Magill SS, et al. New Engl J Med 2014;370:1198

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Central Lines: Utilization

Characteristic	All Patients (N=11,282)	Patients without Health Care-Associated Infections (N=10,830)	Patients with Health Care-Associated Infections (N=452)	P Value†
Central catheter in place on survey date — no. (%)				
Any	2,121 (18.8)	1,862 (17.2)	259 (57.3)	<0.001
Femoral	54 (0.5)	44 (0.4)	10 (2.2)	
Peripherally inserted	1,017 (9.2)	878 (8.1)	159 (35.2)	
Other known type	1,057 (9.4)	958 (8.8)	99 (21.9)	
Unknown type	32 (0.3)	29 (0.3)	3 (0.7)	
None	9,140 (81.0)	8,948 (82.6)	192 (42.5)	
Missing data	21 (0.2)	20 (0.2)	1 (0.2)	

Magill SS, et al. New Engl J Med 2014;370:1198

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Central Line Utilization by Unit

High: ICUs (Medical and Surgical)
Low: Psych, L&D/Postpartum, Ortho

Type of acute care hospital location	No. of locations	Central line days	Patient days	Percentile					
				10%	25%	50% (median)	75%	90%	
Medical/surgical: major teaching	358 (150)	900719	1,482,058	0.54	0.28	0.30	0.53	0.63	0.71
Medical/surgical: all other: <15 beds	1,647 (1,627)	1,260,781	3,473,458	0.37	0.11	0.19	0.34	0.50	0.62
Medical/surgical: all other: >15 beds	807	2,132,226	4,391,341	0.49	0.30	0.40	0.51	0.60	0.69
Neurological	59 (56)	80,894	171,989	0.47	0.22	0.32	0.46	0.55	0.67
Pediatric cardthoracic	43	146,528	202,899	0.72	0.49	0.59	0.75	0.86	0.91
Pediatric medical	31 (29)	23,719	63,991	0.37	0.10	0.14	0.25	0.34	0.47
Pediatric medical/surgical	315 (307)	389,969	866,418	0.45	0.14	0.22	0.35	0.50	0.62
Pediatric surgical	6	3,055	6,009	0.52					
Perinatal	8	710	9,133	0.08					
Psychiatry	6	9,842	26,288	0.37					
Surgical: major teaching	197	470,884	1,133,843	0.57	0.38	0.46	0.57	0.67	0.75
Surgical: all other	199 (198)	245,261	573,964	0.43	0.26	0.31	0.38	0.47	0.54
Surgical: cardthoracic	455 (454)	955,534	1,414,114	0.68	0.45	0.52	0.65	0.74	0.81
Transplant	147	379,688	517,962	0.74	0.50	0.61	0.68	0.77	0.84
Step-down units									
Adult step-down (postcritical care)	700 (699)	818,476	1,714,616	0.48	0.29	0.34	0.42	0.50	0.57
Step-down/ICU (level II)	47 (44)	4,880	57,086	0.31					
Pediatric step-down (postcritical care)	17	17,416	57,086	0.31					
Mixed acute care									
Adult mixed acuity	83 (82)	83,286	336,340	0.25	0.04	0.10	0.19	0.30	0.40
Mixed age mixed acuity	49	28,758	264,837	0.14	0.03	0.06	0.10	0.20	0.32
Pediatric mixed acuity	16	29,340	125,440	0.23					
Long-term acute									
Long-term acute									

$\text{Device utilization ratio} = \frac{\text{No. of device days}}{\text{No. of patient days}}$

Dudeck et al. AJIC 2013; 43: 206-221

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Central Line Associated BSI (CLABSI) Rate by Unit

High: Burn, ICUs (Medical and Surgical), Trauma, Vent Unit
Low: Ortho, GYN, Psych

Type of acute care hospital location	No. of locations	No. of CLABSIs	Central line days	Percentile	Percentile				
					10%	25%	50% (median)	75%	90%
Critical care	73 (69)	219	74,940	2.9	0.0	0.0	2.2	4.4	7.3
Medical: major teaching	251 (250)	812	489,976	1.2	0.0	0.4	1.0	1.8	2.8
Medical: all other	453 (422)	660	2,113,844	1.0	0.0	0.0	0.5	1.4	2.5
Medical: cardiac	387 (381)	583	1,074,844	1.0	0.0	0.0	0.8	1.6	2.6
Medical/surgical: major teaching	749 (744)	696	3,060,679	1.0	0.0	0.0	0.9	1.6	2.6
Medical/surgical: all other: <15 beds	1,847 (1,816)	1,832	1,206,791	0.8	0.0	0.0	0.9	1.9	2.8
Medical/surgical: all other: >15 beds	807 (804)	1,752	2,132,226	0.8	0.0	0.0	0.6	1.2	2.0
Neurological	59 (56)	81	169,896	1.1	0.0	0.0	0.9	1.6	2.8
Neurological	181 (176)	389	387,482	0.9	0.0	0.0	0.7	1.4	2.2

$\text{Device-associated infection rate} = \frac{\text{No. of device-associated infections for an infection site}}{\text{No. of device days}} \times 1,000$

Dudeck et al. AJIC 2015; 43: 206-221

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Definitions: IMPORTANT!


CLINICAL DEFINITION

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SURVEILLANCE DEFINITION

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CLINICAL DEFINITION What is BSI?



Bloodstream infection or Bacteremia:


Positive blood culture(s) +/- systemic signs of infection

Other terms:

- Septicemia: positive blood cultures + systemic signs of infection
- Sepsis and Septic Shock
- Pseudobacteremia or "contaminated" blood cultures: positive blood cultures resulting from contamination during the collection procedure or during laboratory processing

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CLINICAL DEFINITION What is BSI?



Primary BSI: NO identifiable originating source on clinical exam and/or diagnostic testing

Secondary BSI: Identifiable, localized infection at a specific site on clinical exam and/or diagnostic testing

GBS BSI Source in Non-pregnant adults	%
Unknown (Primary)	30-40%
Skin and Soft Tissue	15-40%
Urinary Tract	5-15%
Upper Respiratory Tract	6-12%
Bone and Joint	2-15%
Cardiac/Endocarditis	2-9%
Central Nervous System	<4%

Ex: Group B Streptococcus BSI

Source: UpToDate

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How do pathogens enter the bloodstream?

Many potential points/mechanisms of entry.

Disruption of skin or mucosal barriers:

- Localized infection advances to become systemic (Secondary BSI)
- Skin disruption, scratches, bug bites
- IV drug abuse
- Invasive devices (central venous catheter)
- Invasive procedures (surgical, dental, scopes)

Transient bacteremic episodes may happen all the time but are usually cleared by the liver/spleen

Host considerations

- Implants/prostheses
- Impaired immunity

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CLINICAL DEFINITION **Signs and symptoms**

Systemic: Fever, chills, rigors, fatigue → hypotension, shock
 Respiratory: hyperventilation → respiratory failure
 Neurologic: Confusion → seizure, coma

Secondary: Other localized infection symptoms

Localized infection Fever Septic Shock + Organ Failure

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SIRS criteria (old):
 WBC > 12K or < 4K or > 10% bands
 RR > 20
 HR > 90
 Temp > 38 or < 36C

Image: https://twitter.com/ICPIC_meeting/status/982198791301283842

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Recognize and Treat Sepsis

Patient with suspected infection

qSOFA ≥ 2? (see A) → No → Sepsis still suspected? → No → Monitor clinical condition; reevaluate for possible sepsis if clinically indicated. → Yes → Assess for evidence of organ dysfunction.

SOFA ≥ 2? (see B) → No → Monitor clinical condition; reevaluate for possible sepsis if clinically indicated. → Yes → Sepsis.

Despite adequate fluid resuscitation, 1. vasopressors required to maintain MAP ≥ 65 mm Hg AND 2. serum lactate level > 2 mmol/L? → Yes → Septic shock.

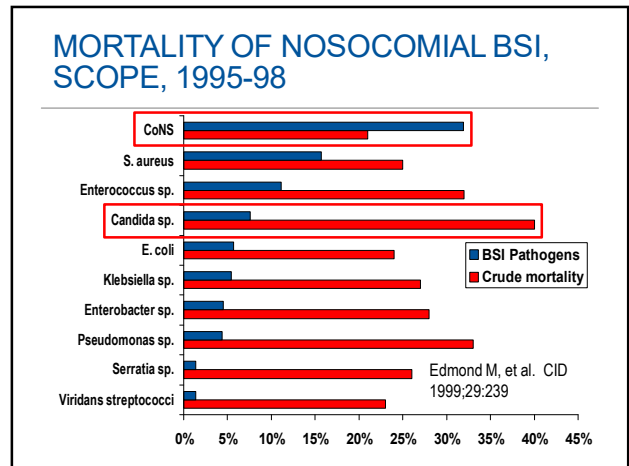
A) qSOFA Variables
 Respiratory rate
 Mental status
 Systolic blood pressure

B) SOFA Variables
 PaO₂/FIO₂ ratio
 Glasgow Coma Scale score
 Mean arterial pressure
 Administration of vasopressors with type and dose rate of infusion
 Serum creatinine or urine output
 Bilirubin
 Platelet count

JAMA. 2016;315(8):801-810

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Clinical management: Go to the Source

- Source control
 - Incision and Drainage for abscesses
 - Remove necrotic material
 - Remove foreign material
 - Contain bowel/bladder contents
 - Wash out joints
- Antibiotics and/or antifungals
 - Initially IV
 - May be able to transition to oral depending on: clinical progress, culture clearance, primary source, and organism/susceptibilities
- Supportive Care
 - Fluids, oxygen, ICU (pressors, vent)

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CLINICAL DEFINITION **Central Venous Catheter Infections**

Infection	Definition
Catheter colonization	Significant growth of ≥1 microorganism in a quantitative or semiquantitative culture of the catheter tip, subcutaneous catheter segment, or catheter hub
Phlebitis	Induration or erythema, warmth, and pain or tenderness along the tract of a catheterized or recently catheterized vein
Exit site infection	
Microbiological	Exudate at catheter exit site yields a microorganism with or without concomitant bloodstream infection
Clinical	Erythema, induration, and/or tenderness within 2 cm of the catheter exit site; may be associated with other signs and symptoms of infection, such as fever or purulent drainage emerging from the exit site, with or without concomitant bloodstream infection*
Tunnel infection	Tenderness, erythema, and/or induration > 2 cm from the catheter exit site, along the subcutaneous tract of a tunneled catheter (e.g., Hickman or Broviac catheter), with or without concomitant bloodstream infection*
Pocket infection	Infected fluid in the subcutaneous pocket of a totally implanted intravascular device; often associated with tenderness, erythema, and/or induration over the pocket, spontaneous rupture and drainage, or necrosis of the overlying skin, with or without concomitant bloodstream infection*
Bloodstream infection	
Infectate related	Concomitant growth of a microorganism from infectate and cultures of percutaneously obtained blood cultures with no other identifiable source of infection
Catheter related	Bacteremia or fungemia in a patient who has an intravascular device and ≥1 positive blood culture result obtained from the peripheral vein, clinical manifestations of infection (e.g., fever, chills, and/or hypotension), and no apparent source for bloodstream infection (with the exception of the catheter). One of the following should be present: a positive result of semiquantitative (≥10 ⁴ cfu per catheter segment) or quantitative (≥10 ⁵ cfu per catheter segment) catheter culture, whereby the same organism (species) is isolated from a catheter segment and a peripheral blood culture; simultaneous quantitative cultures of blood with a ratio of ≥2:1 (cfu/ml) of blood obtained through a peripheral blood; differential time to positivity (growth in a culture of blood obtained through a catheter hub is detected by an automated blood culture system at least 2 h earlier than a culture of simultaneously drawn peripheral blood of equal volume). Note that this definition differs from the definition of central line-associated bloodstream infection used for infection-control surveillance activities.

Mermel L, et al. CID 2009;49:1-45

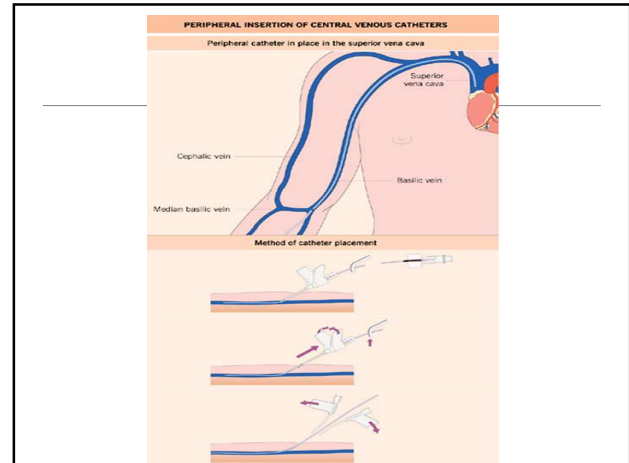
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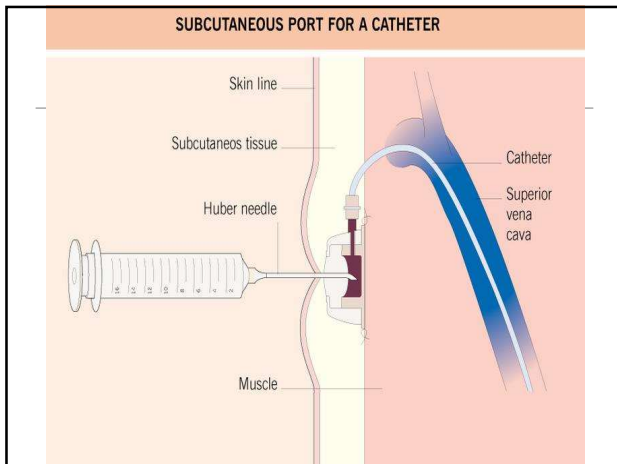
Table 3. Types of intravascular devices and comments on their use.

Type of intravascular device	Comment
Peripheral venous catheter	Usually inserted into the veins of the forearm or the hand; the most commonly used short-term intravascular device
Peripheral arterial catheter	For short-term use; commonly used to monitor hemodynamic status and to determine blood gas levels of critically ill patients; risk of bloodstream infection may approach that of CVCs
Midline catheter	Peripheral catheter (size, 7.6–20.3 cm) is inserted via the antecubital fossa into the proximal basilic or cephalic veins, but it does not enter central veins; it is associated with lower rates of infection, compared with CVCs
Short-term CVC	Most commonly used CVC, accounts for the majority of all catheter-related bloodstream infections
Pulmonary artery catheter	Inserted through a teflon introducer and typically remains in place for an average duration of only 3 days
Pressure-monitoring system	Used in conjunction with arterial catheter; associated with both epidemic and endemic nosocomial bloodstream infections
Peripherally inserted central catheter	Provides an alternative to subclavian or jugular vein catheterization; is inserted via the peripheral vein into the superior vena cava, usually by way of cephalic and basilic veins; similar risk of infection as CVCs in patients hospitalized in intensive care units
Long-term CVC	Surgically implanted CVC (e.g., Hickman, Broviac, or Groshong catheter) with the tunneled portion exiting the skin and a dicron cuff just inside the exit site; used to provide vascular access to patients who require prolonged chemotherapy, home-infusion therapy, or hemodialysis
Totally implantable device	A subcutaneous port or reservoir with self-sealing septum is tunneled beneath the skin and is accessed by a needle through intact skin; associated with low rates of infection

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PATHOGENESIS Central Line Infection

Multifactorial and complex

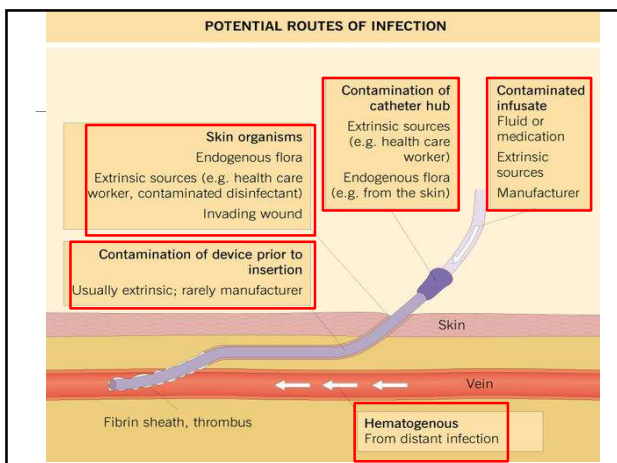
Most catheter-related infections appear to result from **migration of skin organisms** at insertion site into the cutaneous tract with eventual colonization of the catheter tip

Catheter hub also important contributor to intraluminal colonization (especially in long-term catheters)

Less common = hematogenous seeding of catheter tip from distant focus of infection or contaminated infusate

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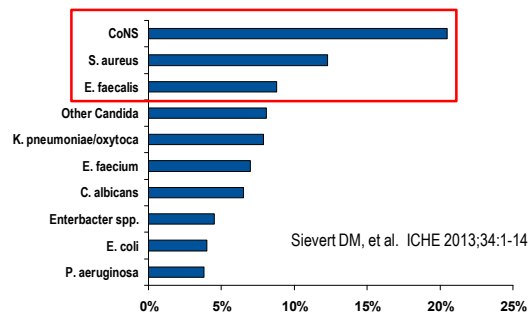
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BIOFILM

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TOP 10 PATHOGENS ASSOCIATED WITH CLABSIs: NHSN, 2009-2010



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Populations at Higher Risk for CLABSI

ICU patients

- High CL utilization (often multiple CL at once and specialized lines)
- Catheters placed in emergencies circumstances
- Need for repeated access daily
- Often need CL for extended time periods

Vulnerable populations

- Hemodialysis
- Peri-operative
- Hem/Onc

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Risk Factors for CLABSI*

- Prolonged hospitalization prior to catheterization
- Prolonged duration of catheterization
- Heavy microbial colonization at the insertion site
- Heavy microbial colonization of the catheter hub
- Site of catheter (adults): Femoral (worst), Internal jugular, compared to Subclavian (best)**
- Multilumen or concurrent catheters
- Substandard catheter care
- Neutropenia
- BMA >40
- Prematurity
- Host Immunity: Neutropenia, neonate prematurity
- Reduced Nurse: Patient Ratios (ICU)
- TPN**
- Substandard catheter care (e.g. excessive manipulation)
- Blood products (children)

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Decreased Risk/Protective Factors

- Female sex
- Antibiotic administration
- Minocycline-rifampin impregnated catheters

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CLINICAL CLUES of CVC INFECTIONS

- CVC: Exit site infection (erythema, tenderness, purulence) or tunnel infection (erythema, tenderness, purulence, induration)
- High grade bacteremia/fungemia (multiple positive cultures)
- Abrupt onset, associated with shock
- Symptoms/signs of sepsis (i.e., fever/ hypotension) without obvious source (no identifiable local infection)
- Evidence of septic thrombophlebitis of great vein
- Continued bacteremia/fungemia despite appropriate therapy
- Symptoms/signs of sepsis plus catheter malfunction
- Bacteremia with CoNS, *Candida*, *Bacillus*, *Corynebacterium*

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COMPLICATIONS OF CLABSIs

Local infection

- Tunnel infection, pocket infection

Sepsis

Remote site infection

- Osteomyelitis
- Meningitis

Endovascular infection

- Endocarditis
- Mycotic aneurysms
- Septic thrombophlebitis

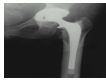
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
Identifying Complicated SAB

Clinical Context Matters


S. aureus Bacteremia + Prosthesis = Trouble




SAB + Arthroplasty = 28% Joint Infection
Murdoch et al *Clin Infect Dis* 2001; 32:647-9.



SAB + Prosthetic Valve = 51% Valve Infection
El-Adhab *Am J Med* 2005; 118:225-9.



SAB + Pacemaker/ICD = 45% Device Infection
Chamis *Circulation* 2001; 104: 1029



SAB + Central Catheter = 71% Thrombophlebitis
Crowley *Crit Care Med* 2008;36:385-90

dicon Slide: Thomas Holland MD, Duke Univ

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Goals of Infection Surveillance: Improve Understanding

Estimate disease incidence:

- Assess program impact, detect outbreaks or problem areas to focus prevention efforts, understand and describe disease burden

Reliability, reproducibility

- Trend over time
- Valid and standardized to the degree possible among practice areas (internal validity)
- Compare to benchmarks (external validity)

The definitions are designed to reflect clinical “truth,” but there is NO method of measurement that is perfect.

Abiding by NHSN definitions improves validity AND provides protection when faced with external review or challenges to the data.

NHSN definitions must be adjusted with time due to the dynamic nature of medicine.

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
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SURVEILLANCE DEFINITION

What is BSI?

There are multiple surveillance definitions to be familiar with for BSI:

- LCBI (1 to 3)
- Secondary BSI due to other site-specific infection
- MBI LCBI (1 to 3)
- CLABSI



https://www.cdc.gov/nhsn/pdfs/pscmanual/4psc_clabscurrent.pdf

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SURVEILLANCE DEFINITION

Healthcare Associated Infection (HAI)

There are multiple “timing” related definitions:

- Date of Event (DOE)
- Healthcare associated infection (HAI)
- Infection window period (IWP)
- Present on admission (POA)
- Repeat infection timeframe (RIT)
- Secondary BSI attribution period (SBAP) = IWP + RIT

An infection is considered an HAI if:

- ALL elements of a CDC/NHSN site-specific infection criterion were **NOT present on admission** but were **ALL present on or after the 3rd calendar day of admission** to the facility.

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SURVEILLANCE DEFINITION

Laboratory Confirmed BSI (LCBI)

Must meet 1 of 3 LCBI criteria:

- Recognized pathogen (1+ cx)
- Common skin commensal (≥2 separate +cx with ≤1 day gap between)
- Neonates + common skin commensal

For all: organism cultured from blood is *not related to infection at another site*

Most closely reflects a clinical Primary BSI.

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
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SURVEILLANCE DEFINITION

LCBI 1

1

Patient of any age has a **recognized pathogen**, which is an organism NOT included on the NHSN common commensal list, identified from **one or more blood specimens obtained** by a culture or non-culture based microbiologic testing method



Recognized Pathogen (Examples)

- S. aureus
- P. aeruginosa
- E. coli
- K. pneumoniae
- S. marcescens
- C. albicans
- Enterococcus sp.

AND

organism cultured from blood is *not related to an infection at another site*

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SURVEILLANCE DEFINITION **LCBI 2**

2

Patient of any age has at least one of the following signs or symptoms: fever (>38.0C), chills, or hypotension

AND

Organism(s) identified from blood is *not related to an infection at another site*

AND

The same NHSN common commensal is identified by a culture or non-culture based microbiologic testing method, from **two or more** blood specimens collected on separate occasions.

Criterion elements must occur within the Infection Window Period (IWP), the 7-day time period which includes the date the positive blood culture was collected, the 3 calendar days before and the 3 calendar days after

Common Commensals (Partial List)

- Diphtheroids [Corynebacterium spp. not C. diphtheriae]
- Bacillus spp. [not B. anthracis]
- Propionibacterium spp.
- Coagulase-negative staphylococci [including S. epidermidis]
- Viridans group streptococci
- Aerococcus spp.
- Micrococcus spp.
- Rhodococcus spp.

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SURVEILLANCE DEFINITION **LCBI 3**

3

Patient **≤ 1 year of age** has at least **one** of the following signs or symptoms: fever (>38.0C), hypothermia (<36.0C), apnea, or bradycardia

AND

Organism(s) identified from blood is *not related to an infection at another site*

AND

The same NHSN common commensal is identified by a culture or non-culture based microbiologic testing method, from **two or more** blood specimens collected on separate occasions.

Criterion elements must occur within the Infection Window Period (IWP), 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.

Common Commensals (Partial List)

- Diphtheroids [Corynebacterium spp. not C. diphtheriae]
- Bacillus spp. [not B. anthracis]
- Propionibacterium spp.
- Coagulase-negative staphylococci [including S. epidermidis]
- Viridans group streptococci
- Aerococcus spp.
- Micrococcus spp.
- Rhodococcus spp.

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SURVEILLANCE DEFINITION **Secondary BSI**

An NHSN site-specific definition must be met; either one of the CDC/NHSN Surveillance Definitions for Specific Types of Infections (defined in Chapter 17), or UTI, PNEU or SSI definitions.

AND

One of the following scenarios must be met:

Scenario 1	Scenario 2
A positive blood specimen must contain at least one eligible matching organism to the site-specific specimen	Positive blood specimen must be an element of the site-specific definition
And the blood specimen is collected in the site-specific secondary BSI attribution period	And blood specimen is collected in the site-specific infection window period
And an eligible organism identified from the site-specific specimen is used as an element to meet the site-specific definition	And an eligible organism identified in a blood specimen is used as an element to meet the site-specific definition

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Table B1: Secondary BSI Guide: List of all NHSN primary site-specific definitions available for making secondary BSI determinations using Scenario 1 or Scenario 2

Scenario 1	Scenario 2
A positive blood specimen must contain at least one eligible matching organism to the site-specific specimen	Positive blood specimen must be an element of the site-specific definition
And the blood specimen is collected in the site-specific secondary BSI attribution period	And blood specimen is collected in the site-specific infection window period
And an eligible organism identified from the site-specific specimen is used as an element to meet the site-specific definition	And an eligible organism identified in a blood specimen is used as an element to meet the site-specific definition

Site	Criterion	Site	Criterion
ABUTL	ABUTL	ABUTL	ABUTL
BONE	1	BONE	1a
BURN	1	BURN	1
CARD	1	DISC	1a
CRIC	2 or 3	ENDO	4a, 4b, 5a or 5b (specific organisms) 6a or 7a plus other criteria as listed
CONJ	1a	ENTD	1
DECU	1	GIT	1a or 2c
DISC	1	HAB	2a or 2b
EAR	1, 3, 5 or 7	JNT	1c
EMRY	1	MEN	2c or 2e
ENDO	1	ORF	1a
EYE	1	PNEU	2 or 3
ISE	2a, 2b (only yeast)	SK	1a
GIT	1 or 2a	UMB	1b
IMB	1	USH	3a or 4b
IC	1		
JHT	1		
LUNG	1		
MED	1		
MEN	1		
ORAL	1, 3a, 3b (only yeast)		
ORF	1		
PJI	1 or 2a		
PNEU	2 or 3		
SK	1		
SKU	1		
SSI	5c, 6b or 6c		
SKN	1a		
ST	1		
UMB	1a		
UR	1a or 2a		
USH	1		
VASC	1a, 1b or 2		
VASC only as SSI	1		
VCGP	1		

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SECONDARY BSI GUIDE FOR ELIGIBLE ORGANISMS (not applicable to VAE)

Figure B1: Secondary BSI Guide for eligible organisms*
(Not applicable to Ventilator-associated Events [VAE], see Figure B2)

```

    graph TD
      A[Positive blood specimen. Site-specific infection suspected as source.] --> B{Is the positive site-specific specimen used as an element to meet the site-specific definition?}
      B -- No --> C{Can the positive blood specimen, which is collected during the infection window period, be used to meet the site-specific infection criterion?}
      C -- No --> D[STOP Primary BSI]
      C -- Yes --> E[Scenario 2 "Element"]
      B -- Yes --> F{Positive blood specimen and site-specific specimen, which is collected during the secondary BSI attribution period, match for at least 1 organism?}
      F -- No --> C
      F -- Yes --> G[Scenario 1 "Match"]
      G --> H[STOP Secondary BSI]
  
```

*Exception: The necrotizing enterocolitis (NEC) definition does not include criteria for a matching site-specific specimen, nor an organism identified from a 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 a 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 but on the same or consecutive days.

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SURVEILLANCE DEFINITION **Laboratory Confirmed BSI (LCBI)**

LCBI Hierarchy: Types of LCBIs (see Table 1 and Table 2):

```

    graph TD
      A[BSIs] --> B[LCBI 1]
      A --> C[LCBI 2]
      A --> D[LCBI 3]
      B --> E[MBI-LCBI 1]
      C --> F[MBI-LCBI 2]
      D --> G[MBI-LCBI 3]
  
```

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Hem/Onc/BMT a “special population” for surveillance

Complex patient population	<ul style="list-style-type: none"> Highly toxic treatments ICU stays Complications (infection, bleeding, ADEs)
Device utilization	<ul style="list-style-type: none"> True need for central line
Culturing practices	<ul style="list-style-type: none"> Bad veins Thrombocytopenia
Antimicrobial utilization	<ul style="list-style-type: none"> Like water Usually appropriate for severity of illness
Surveillance practices	<ul style="list-style-type: none"> Variable?
Administrative pressure	<ul style="list-style-type: none"> “Protective” of program and reputation
Adjudication	<ul style="list-style-type: none"> Clinicians don't consider many “CLABSI” to be preventable Definitions don't apply well to patient population and leads to rejection of data

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SURVEILLANCE
DEFINITION

Mucosal Barrier Injury Laboratory Confirmed Bloodstream Infection (MBI-LCBI)

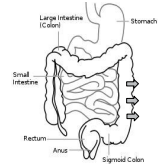
A subset of the LCBI criteria

Aim to identify the “special population” of patients who have BSI due to hematogenous spread from the gut/oral flora due to immune compromise

Must meet one of the 3 LCBI criteria

AND

Must meet one of the two MBI criteria



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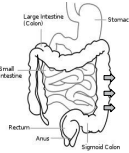
44

SURVEILLANCE
DEFINITION

Mucosal Barrier Injury Laboratory Confirmed Bloodstream Infection (MBI-LCBI)

One of two MBI criteria:

- 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 the 7 calendar days before the date the positive blood specimen was collected.
- Is **neutropenic**, defined as at least two separate days with ANC and/or WBC values < 500 cells/mm³ collected within a 7-day time period which includes the collection date of the positive blood specimen, the 3 calendar days before and the 3 calendar days after.



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SURVEILLANCE
DEFINITION

MBI-LCBI 1

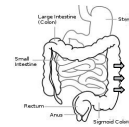
1

LCBI 1 = at least 1 culture positive for “recognized pathogen”

from the intestines

Plus

One of the MBI criteria



Intestinal pathogens
(partial list)

Bacteroides spp.
Candida spp.
Clostridium spp.
Enterococcus spp.
Fusobacterium spp.
Peptostreptococcus spp.
Prevotella spp.
Veillonella spp.
Enterobacteriaceae

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SURVEILLANCE
DEFINITION

MBI-LCBI 2

2

LCBI 2 = signs and symptoms AND at least 2 separate cultures with “common commensals”

Only **viridans group Streptococci** and/or **Rothia** spp. and no other organisms.

Plus

One of the two MBI criteria



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SURVEILLANCE
DEFINITION

MBI-LCBI 3



LCBI 3 = Patient ≤ 1 year of age, AND at least 2 separate cultures with “common commensals”

Only **viridans group streptococci** and/or **Rothia** spp. and no other organisms.

Plus

One of the MBI criteria



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
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SURVEILLANCE DEFINITION

Central Line



Central line: terminates at or close to the heart or in one of the great vessels, used for infusions, withdrawal of blood, or hemodynamic monitoring.

- Catheter must terminate in aorta, pulmonary artery, superior or inferior vena cava, brachiocephalic veins, internal jugular veins, subclavian veins, external iliac or common iliac veins, femoral veins, umbilical artery/vein (neonates)

The following are NOT considered central lines:

- Extracorporeal membrane oxygenation (ECMO)
- Arterial catheters
- Intra-aortic balloon pump (IABP) devices
- Hemodialysis reliable outflow (HeRO) dialysis catheters
- Non-accessed central line (not accessed nor inserted during the hospitalization)
- Peripheral IV or Midlines
- Ventricular Assist Device (VAD)

Infusion: Introduction of a solution through a blood vessel via a catheter lumen

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SURVEILLANCE DEFINITION

Central Line: Temporary, Permanent

Temporary central line: A non-tunneled, non-implanted catheter

Permanent central line:

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

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

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SURVEILLANCE DEFINITION

CENTRAL LINE-ASSOCIATED BLOODSTREAM INFECTION (CLABSI) EVENT

Eligible Central Line: A CL that has been in place for **more than two** consecutive calendar days (on or after CL day 3), following the **first access** of the central line, in an inpatient location, during the current admission. Such lines are eligible for CLABSI events and remain eligible for CLABSI events **until the day after removal from the body or patient discharge**, whichever comes first.

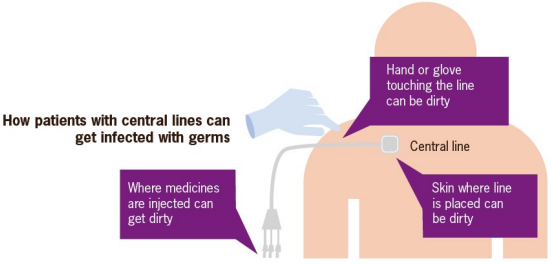
Central line-associated BSI (CLABSI): A laboratory confirmed bloodstream infection (LCBI) where an eligible BSI organism is identified and an eligible central line is present on the LCBI date of event or the day before.

https://www.cdc.gov/nhsn/pdfs/pscmanual/4psc_clabscurrent.pdf

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How does CLABSI happen?



How patients with central lines can get infected with germs

CDC VitalSigns March 2011,60(8):243-248.

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Contamination occurs...

Insertion:

- Patient's Skin
- Operator (Spit, Hair, Hands)
- Environment

Maintenance:

- Cap is frequently accessed, inadequately cleaned during access, or poorly functioning
- Operator (Spit, Hair, Hands) during assessments + routine dressing changes
- Bacterial migration along catheter tract from skin

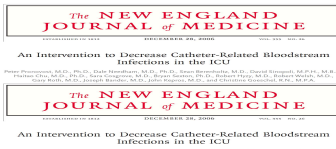
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What's a Bundle?

I Institute for Healthcare Improvement

"Care bundles are small, straightforward, sets of evidence-based practices... that, when implemented collectively, improve the reliability of their delivery and improve patient outcomes."



<http://www.ihl.org/resources/Pages/ImprovementStories/WhatsaBundle.aspx>

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

IHI Bundle: PREVENTION OF CENTRAL LINE INFECTIONS

During insertion:

- Hand hygiene
- Maximal barrier precautions
- Chlorhexidine skin antiseptis (now CHG-alcohol)
- Optimal catheter site selection, with subclavian vein as the preferred site for nontunneled catheters


During maintenance:

- Daily review of line necessity, with prompt removal of unnecessary lines

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
Infection Control & Hospital Epidemiology (2022), 1-17
doi:10.1017/ice.2022.87



SHEA/IDSA/APIC Practice Recommendation

Strategies to prevent central line-associated bloodstream infections in acute-care hospitals: 2022 Update

Niccolò Buetti MD, MSc, PhD^{1,2,9}, Jonas Marshall MD, MSc^{4,8}, Marci Drees MD, MS^{5,6}, Mohamad G. Fakih MD, MPH⁷, Lynn Hadaway MEd, RN, NPD-BC, CRNP⁸, Lisa L. Maragakis MD, MPH⁸, Elizabeth Monsees PhD, MBA, RN, CIC^{10,11}, Shannon Novosad MD MPH¹², Naomi P. O'Grady MD¹³, Mark E. Rupp MD¹⁴, Joshua Wolf MBBS, PhD, FRACP^{15,16}, Deborah Yokoe MD, MPH¹⁷ and Leonard A. Mermel DO, ScM^{18,19}




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
GRADING THE QUALITY OF EVIDENCE

Category	Definition
HIGH	Highly confident that the true effect lies close to that of the estimated size and direction of the effect. Evidence is rated as high quality when there are a wide range of studies with no major limitations, there is little variation between studies, and the summary estimate has a narrow confidence interval.
MODERATE	The true effect is likely to be close to the estimated size and direction of the effect, but there is a possibility that it is substantially different. Evidence is rated as moderate quality when there are only a few studies and some have limitations but not major flaws, there is some variation between studies, and/or the confidence interval of the summary estimate is wide.
LOW	The true effect may be substantially different from the estimated size and direction of the effect. Evidence is rated as low quality when supporting studies have major flaws, there is important variation between studies, the confidence interval of the summary estimate is very wide, and/or there are no rigorous studies.

*Based on the CDC Healthcare Infection Control Practices Advisory Committee (HICPAC) "Update to the Centers for Disease Control and Prevention and the Healthcare Infection Control Practice Advisory Committee Recommendations: Categorization Scheme for Infection Control and Prevention Guideline Recommendations" (October 2019), the Grades of Recommendation, Assessment, Development, and Evaluation (GRADE),¹⁰ and the Canadian Task Force on Preventive Health Care.¹¹



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Essential Practices

Before insertion

- Provide easy access to an evidence-based list of indications for CVC use to minimize unnecessary CVC placement. (Quality of Evidence: LOW)
- Require education and competency assessment of HCP involved in insertion, care, and maintenance of CVCs about CLABSI prevention. (Quality of Evidence: MODERATE)^{10,11}
- Bathe ICU patients aged >2 months with a chlorhexidine preparation on a daily basis. (Quality of Evidence: HIGH)^{10,11}

At insertion

- In ICU and non-ICU settings, a facility should have a process in place, such as a checklist, to ensure adherence to infection prevention practices at the time of CVC insertion. (Quality of Evidence: MODERATE)^{10,11}
- Perform hand hygiene prior to catheter insertion or manipulation. (Quality of Evidence: MODERATE)^{10,11,12}
- The solution should be preferred to reduce infection complications when the catheter is placed in the ICU setting. (Quality of Evidence: HIGH)^{10,11,13,14}
- Use an all-inclusive catheter cap or AI. (Quality of Evidence: MODERATE)^{10,11}
- Use an antiseptic barrier for catheter insertion. (Quality of Evidence: MODERATE)^{10,11}
- Use maximum sterile barrier precautions during CVC insertion. (Quality of Evidence: MODERATE)^{10,11,15}
- Use an alcoholic chlorhexidine antiseptic for skin preparation. (Quality of Evidence: HIGH)^{10,11,16}

After insertion

- Ensure appropriate nurse-to-patient ratio and limit use of float nurses in ICU. (Quality of Evidence: HIGH)^{10,17}
- Use chlorhexidine-containing dressings for CVCs in patients over 2 months of age. (Quality of Evidence: HIGH)^{10,11,18,19}
- For non-tunneled CVCs in adults and children, change transparent dressings and perform site care with a chlorhexidine-based antiseptic at least every 7 days or immediately if the dressing is soiled, loose, or damp. Change gauze dressings every 2 days or earlier if the dressing is soiled, loose, or damp. (Quality of Evidence: MODERATE)^{10,11,20}
- Disinfect catheter hubs, medline connectors, and injection ports before accessing the catheter. (Quality of Evidence: MODERATE)^{10,21,22}
- Remove unnecessary catheters. (Quality of Evidence: MODERATE)^{10,23}
- Routine replacement of administration sets not used for blood, blood products, or lipid formulations can be performed at intervals up to 7 days. (Quality of Evidence: HIGH)^{10,24}
- Perform surveillance for CLABSI in ICU and non-ICU settings. (Quality of Evidence: HIGH)^{10,25,26}

Additional Practices


- Use antibiotic- or antimicrobial-impregnated CVCs. (Quality of Evidence: HIGH in adult patients)^{10,27,28,29} and Quality of Evidence: MODERATE in pediatric patients.^{10,30}
- Use antimicrobial lock therapy for long-term CVCs. (Quality of Evidence: HIGH)^{10,31,32}
- Use recombinant tissue plasminogen activating factor (rTPA) once weekly after hemodialysis in patients undergoing hemodialysis through a CVC. (Quality of Evidence: HIGH)^{10,33}
- Utilize infrared or vascular access beams for reducing CLABSI rates. (Quality of Evidence: LOW)^{10,34}
- Use antimicrobial cement for hemodialysis catheter insertion sites. (Quality of Evidence: HIGH)^{10,35}
- Use an antibiotic-containing hub/connector cap/port protector to cover connectors. (Quality of Evidence: MODERATE)^{10,36,38}

Practices that should not be considered a routine part of CLABSI prevention

- Do not use antimicrobial prophylaxis for short-term or tunneled catheter insertion or while catheters are in situ. (Quality of Evidence: HIGH)^{10,37,38}
- Do not routinely replace CVCs or arterial catheters. (Quality of Evidence: HIGH)¹⁰

Unintended consequences

- Routine use of needles/connectors as a CLABSI prevention strategy before an assessment of risks, benefits, and education regarding proper use.^{10,39}
- Surveillance of other types of catheters (eg, peripheral arterial or peripheral venous catheters).^{10,40}
- Standardized, nonantimicrobial management dressing and CLABSI risk.
- The impact of using chlorhexidine-based products on bacterial resistance to chlorhexidine.
- Suboptimal assessment.
- Impact of other antiseptic-impregnated umbilical catheters in preterm infants (especially in countries where it is approved for use in children).¹⁰
- Necessity of mechanical disinfection of a catheter hub, medline connector, and injection port before accessing the catheter when antiseptic-containing caps are being used.




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PREVENTING CLABSI: BEFORE INSERTION

Provide easy access to an evidence-based list of indications for CVC {Low}

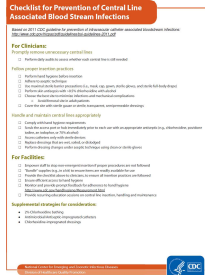

Require education of HCP involved in insertion, care, and maintenance of CVCs about CLABSI prevention {Moderate}

Bathe ICU patients over 2 mo of age with a CHG preparation on a daily basis {High}




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CDC EDUCATIONAL MATERIAL

<http://www.cdc.gov/HAI/bsi.html>



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BATHE ICU PATIENTS >2 MONTHS OF AGE WITH A CHG PREPARATION DAILY

Intervention = Daily bathing with 2% CHG impregnated washcloth
 Design & setting : Cross-over study in MICU
 Result: CHG associated with decreased rate (per 1,000 pt-days) of CLABSI (4.1 vs 10.4)

Washcloth period: 28 weeks (July 8-October 20, 2005) and 24 weeks (January 5-June 21, 2006).
 Group A: 2% Chlorhexidine cloths and Soap and water.
 Group B: Soap and water and 2% Chlorhexidine cloths.

Bleasdale S, et al. Arch Intern Med 2007;167:2073

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REDUCE MRSA

Cluster-randomized trial in 74 ICUs
 MRSA screening and isolation vs. targeted decolonization of MRSA carriers vs. universal decolonization
 Decolonization: CHG daily bathing + nasal mupirocin
 Result: Universal decolonization reduced rate of all Primary BSI significantly. Decreased MRSA BSI also, but NS.
 1 BSI prevented per 99 patients decolonized.

Huang SS et al. N Engl J Med 2013;368:2255-2265.
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PREVENTING CLABSI: AT INSERTION

- Have a process in place to ensure adherence to infection prevention practices (e.g., checklist) (Moderate)
- Perform hand hygiene prior to catheter insertion or manipulation (Moderate)
- Subclavian site is preferred in the ICU setting (Avoid using the femoral artery for central venous access in obese patients) (High)
 - Consider risks and benefits of different insertion sites
 - Do not use peripherally inserted CVCs (PICCs) as a strategy to reduce CLABSI
- Use an all-inclusive catheter cart or kit (Moderate)
- Use ultrasound guidance for internal jugular insertion (High)
- Use maximum sterile barrier precautions during CVC insertion (mask, cap, sterile gown, and sterile gloves; patient covered with full body sterile drape) (Moderate)
- Use alcohol-chlorhexidine for skin antisepsis (High)

CVC Bundle Checklist:

- Hand Hygiene
- Mask, cap, gown, sterile gloves, full body drape
- CHG-alcohol skin antisepsis
- Optimal line site selection

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CLABSI: MAINTENANCE

- Ensure appropriate nurse-to-patient ratio and limit the use of float nurses in the ICU (High)
- Use CHG-containing dressings for CVCs in patients > 2 months (High)
- Disinfect catheter hubs, needleless connectors, and injection ports before accessing the catheter (Moderate)
- Remove nonessential catheters (Moderate)
- For non-tunneled CVCs, change dressings and perform site care with CHG-based antiseptic q7d or pm damp/loose/soiled dressing. Gauze q2 days or pm damp/loose/soiled dressing (Moderate)
- Replace administration sets not used for blood, blood products, or lipids at intervals not longer than 96 hours (High)

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PREVENTING CLABSI: SPECIAL APPROACHES

- Use antiseptic or antimicrobial-impregnated CVCs in adult patients (High/Moderate) in specific situations:
 - Higher than desired CLABSI rate
 - Patients with recurrent CLABSI
 - Patients at higher risk of severe sequelae from a CLABSI (e.g. prosthetic valves)
- Use an antiseptic-containing hub/connector cap/port protector to cover connectors (Moderate)
- Use recombinant tPA for HD through CVC (High)
- Use vascular access teams (Low)
- Use antimicrobial locks for CVCs (High) in specific situations:
 - HD catheters
 - Limited access and history of recurrent CLABSI
 - Patients at higher risk of severe sequelae from a CLABSI
- AVOID:
 - Antimicrobial prophylaxis
 - Routine replacement of CVCs

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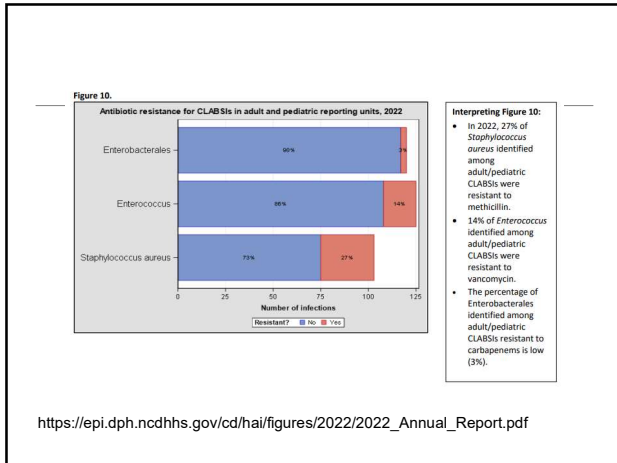
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PREVENTING CLABSI: UNRESOLVED ISSUES

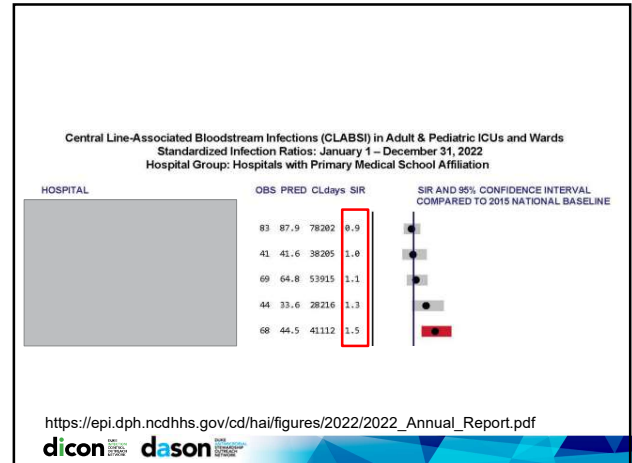
- Routine use of needleless connectors
- Silver-coated catheters
- Standard transparent dressings (nonantimicrobial)
- Impact of CHG-containing products on CHG-resistance
- Sutureless securement
- Necessity of manual disinfection of hub/needleless connector when antiseptic-caps used

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IC effect on primary BSI

The Effect of a Nationwide Infection Control Program Expansion on Hospital-Onset Gram-Negative Rod Bacteremia in 130 Veterans Health Administration Medical Centers: An Interrupted Time-Series Analysis

Horizontal

- Local MDRO coordinator
- Culture transformation
- Education
- Leadership

Vertical (MRSA+ only)

- Active surveillance
- Contact precautions

Figure 1. Effect of the Methicillin-Resistant *Staphylococcus aureus* Prevention Initiative on changes in incidence rates of gram-negative rod bacteremia. Solid slope lines are slopes estimated by autoregressive models; break slope lines are estimated slopes without effects of intervention; vertical break lines are beginning and end of implementation of the initiative.

CID. 2016; 63 (5):642-50.

<https://www.cdc.gov/hai/data/portal/progress-report.html>

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CLABSI Prevention Success!

In 2017, there were 24,265 CLABSIs reported by 3576 United States acute care hospitals to the United States Centers for Disease Control and Prevention's National Healthcare Safety Network

-19%

Prevention efforts have saved ~ 3,000-6,000 lives and ~\$414 million in extra medical costs (2009 compared with 2001)

United States Centers for Disease Control and Prevention. Current HAI Progress Report. <https://www.cdc.gov/hai/data/portal/progress-report.html> MMWR Morb Mortal Wkly Rep. 2011;60(8):243.

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But, CLABSIs still occur: ~30,000 per year

Nationally, among acute care hospitals, the 2021 annual highlights in this report include:

- Overall, 7% increase in CLABSI between 2020 and 2021
 - Largest increase in ICUs (10%)

<https://www.cdc.gov/hai/data/portal/progress-report.html>

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CONCLUSIONS

- Healthcare-associated bloodstream (BSI) cause significant morbidity and mortality
- The most important risk factor for BSI is presence of a central venous catheter
- Clinical definition and surveillance definition of catheter-related BSI are NOT the same
- A near 0 rate of CLABSI is possible using existing technology and appropriate practice strategies
- Current guidelines should be followed for the prevention of CLABSI

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Key References

Clinical Management of catheter-related infections.

▪ *Clinical Infectious Diseases*; 2009; 49: 1-45.

Prevention of catheter-related infections.

▪ *Clinical Infectious Diseases*; 2011; 52: e1-e32.

SHEA Compendium: Strategies to Prevent CLABSI.

▪ *Infection Control & Hospital Epidemiology* (2022), 1–17

Sepsis-3 definition and management.

▪ *JAMA*. 2016;315(8):801-810.