Central Line-Associated Bloodstream Infection (CLABSI) Surveillance in Oncology and Bone Marrow/Solid Organ Transplant

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CME Disclosure

Dr. Srigley does not report any conflict of interest and does not intend to make therapeutic recommendations for medications that have not received regulatory approval (i.e. “off-label” use of medication).
Objectives

To understand the importance of CLABSI surveillance in oncology/bone marrow transplant (BMT)/solid organ transplant (SOT) settings

To review the surveillance definitions of CLABSI

To discuss best practices for CLABSI prevention and surveillance
Importance of CLABSI Surveillance in Oncology/BMT/SOT
Burden of CLABSIs in Oncology

BELLONI ET AL, WORLDV EVID-BASED NU 2022;19:100–111

BAIER ET AL, PLOS ONE 2020;15(1): E0227772

10.6 cases per 10,000 CVC days
Impacts of CLABSI

Patient outcomes

Study in hematology-oncology population found non-statistically significant increase in mortality (7% vs. 4%, p = 0.115)

Systematic review found overall odds ratio of in-hospital death associated with CLABSI = 2.75 (CI 1.86–4.07)

Length of stay was double for patients with CLABSI (30 vs. 15 days, p<0.01)

Financial costs

Median costs attributable to CLABSI = EUR 8,810

Ziegler et al, Infection 2015;43:29-36
CLABSI Is Preventable

Overall incidence rate ratio of 0.459 (95% CI, 0.381-0.554)
Changes in CLABSI rates up to 100%
CLABSI has the highest number of preventable deaths and the highest cost impact of any HAI

Schreiber et al, Infect Control Hosp Epidemiol 2018;39:1277-95
CLABSI Definitions
Laboratory-Confirmed Bloodstream Infection (LCBI)

1) Recognized bacterial or fungal pathogen, not included on the NHSN common commensal list, identified from one or more blood specimens by culture or non-culture based microbiologic testing methods

2) The same NHSN common commensal is identified by a culture from two or more blood specimens collected on separate occasions AND patient has at least one of the following signs or symptoms: fever (>38.0°C), chills, or hypotension

3) The same NHSN common commensal is identified by a culture from two or more blood specimens collected on separate occasions AND patient ≤ 1 year of age has at least one of the following signs or symptoms: fever (>38.0°C), hypothermia (<36.0°C), apnea, or bradycardia

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

- Actinomyces spp.
- Aerococcus spp.
- Bacillus spp.
- Corynebacterium spp.
- Cutibacterium acnes
- Diphtheroids
- Kocuria spp.
- Micrococcus spp.
- Paenibacillus spp.
- Propionibacterium spp.
- Rothia spp.
- Coagulase-negative staphylococci
- Viridans group streptococci

http://www.cdc.gov/nhsn/xls/master-organism-com-commensals-lists.xlsx
Primary vs. Secondary BSI

Primary: LCBI that is not secondary to an infection at another body site

Secondary: BSI thought to be seeded from a site-specific infection

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

Primary LCBI where an eligible central line is present on the day of LCBI diagnosis or the day before

CL must have been in place for more than 2 consecutive calendar days following the first access of the central line, in an inpatient location, during the current admission

Types of CL

Permanent central line, which includes:
- Tunneled catheters, including tunneled dialysis catheters
- Implanted catheters (including ports)

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

Umbilical catheter: A vascular catheter inserted through the umbilical artery or vein in a neonate. All umbilical catheters are central lines

https://www.cdc.gov/nhsn/pdfs/pscmanual/4psc_clabscurrent.pdf
Mucosal Barrier Injury LCBI

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:

a. Grade III or IV gastrointestinal graft versus host disease [GI GVHD] OR
b. ≥1-liter diarrhea in a 24-hour period (or ≥20 mL/kg in a 24-hour period for patients <18 y/o) with onset within the 7 calendar days before collection date of the positive blood specimen OR

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

AND

LCB1 + at least one blood specimen with ONLY intestinal organisms from the NHSN MBI organism list, OR
LCBI 2 or 3 + at least 2 matching blood specimens with ONLY viridans group Streptococcus and/or Rothia spp. alone but no other organisms

https://www.cdc.gov/nhsn/pdfs/pscmanual/4psc_clabscurrent.pdf
Best Practices for CLABSI Prevention and Surveillance
SHEA/IDSA/APIC Practice Recommendation

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

Niccolò Buetti MD, MSc, PhD, Jonas Marschall MD, MSc, Marci Drees MD, MS, Mohamad G. Fakih MD, MPH, Lynn Hadaway MEd, RN, NPD-BC, CRNI, Lisa L. Maragakis MD, MPH, Elizabeth Monsees PhD, MBA, RN, CIC, Shannon Novosad MD MPH, Naomi P. O’Grady MD, Mark E. Rupp MD, Joshua Wolf MBBS, PhD, FRACP, Deborah Yokoe MD, MPH and Leonard A. Mermel DO, ScM
Before Insertion

List of indications for CVC use to avoid unnecessary CVC placement

Education and competency assessment of HCP involved in insertion, care, and maintenance of CVCs

Daily chlorhexidine bathing (ICU patients aged >2 months)

Some studies suggest benefit among adult hematology-oncology patients but not for pediatric patients; need to consider risks (e.g. increased resistance) and costs

Buetti et al, Infect Control Hosp Epidemiol 2022;43:553-69
At Insertion

Checklist to ensure adherence to CLABSI prevention practices
Hand hygiene prior to insertion or manipulation
Subclavian site is preferred
All-inclusive catheter cart or kit
Ultrasound guidance for insertion
Maximum sterile barrier precautions
Alcoholic chlorhexidine antiseptic for skin preparation

Buetti et al, Infect Control Hosp Epidemiol 2022;43:553-69
After Insertion

Ensure appropriate nurse-to-patient ratio and limit use of float nurses in ICUs

Chlorhexidine-containing dressings for CVCs

Change transparent dressings and perform site care with chlorhexidine at least q 7 days; change gauze dressings q 2 days

Disinfect catheter hubs, needleless connectors, and injection ports before access

Remove nonessential catheters

Routine replacement of administration sets can be performed at intervals up to 7 days

Perform surveillance for CLABSI

Buetti et al, Infect Control Hosp Epidemiol 2022;43:553-69
Additional Approaches

- Use antiseptic- or antimicrobial-impregnated CVCs
- Use antimicrobial lock therapy for long-term CVCs
- Utilize infusion or vascular access teams for reducing CLABSI rates
- Use an antiseptic-containing hub/connector cap/port protector to cover connectors
- Do not use antimicrobial prophylaxis for short-term or tunneled catheter insertion or while catheters are in situ
- Do not routinely replace CVCs or arterial catheters

Buetti et al, Infect Control Hosp Epidemiol 2022;43:553-69
Surveillance Best Practices

Use consistent methods and definitions to allow comparison to benchmark data

Denominator data collection

Methods

- Manual, daily
- Manual, once/week (not recommended for oncology/specialty care areas)
- Electronic

For oncology/specialty care areas, separate counts for permanent vs temporary CLs
Only 1 CL per patient is counted per day regardless of # of CLs present
All CLs should be included in device day counts regardless of access

Buetti et al, Infect Control Hosp Epidemiol 2022;43:553-69
https://www.cdc.gov/nhsn/pdfs/pscmanual/4psc_clabscurrent.pdf
Guidelines for Preventing Infectious Complications among Hematopoietic Cell Transplantation Recipients: A Global Perspective

Marcie Tomblyn, Tom Chiller, Herrmann Einsele, Ronald Gress, Kent Sepkowitz, Jan Storek, John R. Wingard, Jo-Anne H. Young, Michael A. Boeckh

MBI-LCBI Prevention

Standard CLABSI prevention bundles

Oral care bundles
  E.g. sodium bicarbonate mouthwash, artificial saliva, lip balm, scheduled nursing oral assessments, dental hygiene

Maintaining a healthy orogastrointestinal microbiome?
  E.g. antimicrobial stewardship, probiotics, dietary interventions

Vaughan et al, Infect Control Hosp Epidemiol 2017;38:1385-7
Dandoy et al, Bone Marrow Transplant 2019;54:1932-9
Kemp et al, J Ped Hematol/Oncol Nurs 2019;36:321-6
Reed et al, JCO Oncol Practice 2020;16:e306-12
Questions?

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