

Appendix: Recommendations for Prevention and Control of Infections in Neonatal Intensive Care Unit Patients: Central Line-associated Blood Stream Infections

Table of Contents

| A. Search Strategies and Results | 5 |
|--|----|
| A.1. Guideline Search Strategies (April 2011) | |
| A.2. Primary Study Search Strategies: Central Line-associated Bloodstream Infections (CLABSI) (May 5, 2021) | |
| A.3. Primary Study Search Strategies: Central Line-associated Bloodstream Infections and Chlorhexidine (May 5, 2021) | 7 |
| B. Study Exclusion Criteria | 9 |
| C. Evidence Review | |
| C.1. Non-sterile Gloves | |
| C.2. Central Line Type | 14 |
| C.3. Central Line Insertion Site | |
| C.4. Number of Catheter Lumens | 47 |
| C.5. Skin Antisepsis for Catheter Insertion and Maintenance | 52 |
| C.6. Chlorhexidine Bathing | 55 |
| C.7. Catheter Hub Manipulation | |
| C.8. Central Line Antimicrobial Locks | 65 |
| C.9. Optimal Umbilical Arterial and Venous Catheter Dwell Time | 71 |
| C.10. Optimal Peripherally Inserted Central Catheter Dwell Time | |
| C 11 Dedicated Catheter Care Team | 94 |

| C.12. Central Line Insertion and Maintenance Bundles | 98 |
|---|--------------------------------|
| C.13. Prophylactic Antimicrobial Administration | 110 |
| C.14. Prophylactic Anticoagulant Administration | 118 |
| D. Evaluation of Study-level Risk of Bias | 127 |
| D.1. Randomized Controlled Trial Checklist | 127 |
| D.2. Observational Study Checklist | 127 |
| D.3. Descriptive Study Checklist | 127 |
| D.4. Rating for Overall Risk of Bias | Error! Bookmark not defined |
| D.4. Aggregate Risk of Bias | 128 |
| E. HICPAC Recommendation Categorization Scheme (2019) | 128 |
| F. References | 131 |
| G. Acronyms and Abbreviations | 136 |
| Tables | |
| Table 1 Guideline Search of MEDLINE | |
| Table 2 Guideline Search of American Academy of Pediatrics (AAP) | |
| Table 3 Primary Search of MEDLINE: CLABSI | 5 |
| Table 4 Primary Search of EMBASE: CLABSI | 6 |
| Table 5 Primary Search of Cochrane Library: CLABSI | 6 |
| Table 6 Primary Search of CINAHL: CLABSI | 7 |
| Table 7 CLABSI and Chlorhexidine Search Strategy for MEDLINE | 7 |
| Table 8 Primary Search of EMBASE: CLABSI and Chlorhexidine | |
| Table 9 Search of the Cochrane Library: CLABSI and Chlorhexidine | |
| Table 10 The Summary of Evidence for Using Non-Sterile Gloves After Hand Hygiene vs. Hand Hygiene Alone | Prior to Every Patient Contact |
| to Prevent CLABSI | 11 |
| Table 11 Extracted Information for Non-Sterile Gloves After Hand Hygiene to Prevent CLABSI | 11 |
| Table 12 Risk of Bias of Randomized Controlled Trials on Using Non-Sterile Gloves After Hand Hygiene | |
| Table 13 The Summary of Evidence on UVC vs. Peripheral Catheters to Prevent CLABSI | 14 |
| Table 14 The Summary of Evidence for the Efficacy of All Catheter Types to Prevent CLABSI | |
| Table 15 Extracted Information on Central Line Type | |
| | |

| Table 16 Risk of Bias of Two Group Studies on Catheter Types | 30 |
|---|-------|
| Table 17 Summary of Findings on Central Line Sites to Prevent CLABSI: PICC Placement in Femoral vs. Non-Femoral Sites | 30 |
| Table 18 Summary of Findings on Central Line Sites to Prevent CLABSI: CVC Placement in Jugular vs. Subclavian vs. Femoral Sites | 31 |
| Table 19 Summary of Findings on the Efficacy of Central Line Site to Prevent CLABSI: CVC Placement in Upper vs. Lower Extremities | 31 |
| Table 20 Extracted Information on Central Line Sites | 32 |
| Table 21 Risk of Bias of Two Group Studies on Catheter Sites | |
| Table 22 Summary of Findings on the Number of Umbilical Venous Catheter Lumens to Prevent CLABSI | 47 |
| Table 23 Extracted Information on the Number of Umbilical Venous Catheter Lumens | |
| Table 24 Risk of Bias for Randomized Controlled Trials on Number of Catheter Lumens | 48 |
| Table 25 Risk of Bias for two group studies on Number of Catheter Lumens | |
| Table 26 Summary of Findings on the Use of 2% alcoholic CHG vs. 10% PI for catheter insertion and maintenance | 52 |
| Table 27 Extracted Information on the Use of Chlorhexidine Skin Antiseptic | 53 |
| Table 28 Risk of Bias for Randomized Controlled Trials Using Chlorhexidine Skin Antiseptics | 54 |
| Table 29 Summary of Findings on Bathing with 2% CHG cloths vs. Placebo or No Bathing to Prevent CLABSI | 55 |
| Table 30 Summary of Findings on a Single Bath with 0.25% CHX Cloths vs. Saline Impregnated Cloths vs. No Cleansing to Prevent CLABS | 31 56 |
| Table 31 Extracted Information on Chlorhexidine Bathing | 56 |
| Table 32 Risk of Bias of Randomized Controlled Trials on Chlorhexidine Bathing | 62 |
| Table 33 Risk of Bias of Two Group Studies on Chlorhexidine Bathing | |
| Table 34 Summary of Findings on Catheter Manipulation to Prevent CLABSI in NICU Patients | |
| Table 35 Extracted Information on Catheter Manipulation | |
| Table 36 Risk of Bias for Two Group Studies on Catheter Hub Manipulation | |
| Table 37 Summary of Findings on Antimicrobial locks vs. Standard of Care to Prevent CLABSI | 66 |
| Table 38 Extracted Information on Central Line Antimicrobial Locks | |
| Table 39 Risk of Bias for Randomized Controlled Trials on Central Line Antimicrobial Locks | |
| Table 40 Summary of Findings on the Optimal Duration of Umbilical Catheters Prior to Prevent CLABSI | 71 |
| Table 41 Summary of Findings on the Optimal Duration of Umbilical Artery Catheter for Removal to Prevent CLABSI | 72 |
| Table 42 Summary of Findings on the Optimal Duration Prior to Removal of Umbilical Venous Catheters to Prevent CLABSI | 73 |
| Table 43 Summary of Findings on the Optimal Duration Umbilical Venous Catheter for Replacement with a Long-term Catheter to Prev | ent |
| CLABSI | |
| Table 44 Extracted Information on Umbilical Catheter Dwell Time | 74 |
| Table 45 Risk of Bias for Randomized Controlled Trials on Umbilical Catheter Dwell Times | |
| Table 46 Risk of Bias for Two Group Studies on Umbilical Catheter Dwell Times | |
| Table 47 Risk of Bias for Single Group Studies on Umbilical Catheter Dwell Times | |
| Table 48 Risk of Bias for Two Group Studies on Umbilical Catheter Dwell Times | 82 |

| Table 49 Summary of Findings on Peripherally Inserted Central Catheter Dwell Times to Prevent CLABSI | 83 |
|---|------|
| Table 50 Extracted Information on Peripherally Central Catheter Dwell Time | 83 |
| Table 51 Risk of Bias for Two Group Studies on Percutaneous Central Catheter Dwell Times | 93 |
| Table 52 Risk of Bias for Single Group Studies on Percutaneous Central Catheter Dwell Times | 93 |
| Table 53 Summary of Findings for a Dedicated Percutaneous Inserted Central Catheter Care Team vs. Standard of Care to Prevent CLABS | ı 94 |
| Table 54 Extracted Information on a Dedicated Percutaneous Inserted Central Catheter Care Team | 94 |
| Table 55 Risk of Bias for Two Group Studies on a Dedicated Percutaneous Inserted Central Catheter Care Team | 98 |
| Table 56 Summary of Findings on Insertion and Maintenance Bundles vs. Standard of Care to Prevent CLABSI | 98 |
| Table 57 Extracted Information for Central Venous Catheter Insertion and Maintenance Bundles | 98 |
| Table 58 Risk of Bias for Two Group Studies on Central Venous Catheter Insertion and Maintenance Bundles | 110 |
| Table 59 Summary of Findings on Prophylactic Amoxicillin vs. No Prophylactic Amoxicillin to Prevent CLABSI | 110 |
| Table 60 Summary of Findings on Prophylactic Vancomycin vs. No Prophylactic Vancomycin to Prevent CLABSI | 110 |
| Table 61 Extracted Information on Prophylactic Antimicrobials | 111 |
| Table 62 Risk of Bias for Randomized Controlled Trials on Prophylactic Antimicrobials | 117 |
| Table 63 Risk of Bias for Two Group Studies on Prophylactic Antimicrobials | 118 |
| Table 64 Summary of Findings on Prophylactic Heparin + TPN or dextrose vs. TPN or dextrose to Prevent CLABSI | 118 |
| Table 65 Extracted Information on Anticoagulant Infusion | |
| Table 66 Risk of Bias for Randomized Controlled Trials on Anticoagulant Infusion | 126 |
| Table 67 Strength of Recommendations | 128 |
| Table 68 Justification for Choice of Recommendation Strength | 129 |
| Table 69 Aggregate Level of Confidence in Effect Estimate* | 130 |

A. Search Strategies and Results

A.1. Guideline Search Strategies (April 2011)

Table 1 Guideline Search of MEDLINE

| # | Search History | Results |
|---|-------------------|---------|
| 1 | As outlined below | 61 |

Table 2 Guideline Search of American Academy of Pediatrics (AAP)

| # | Search History | Results |
|---|------------------------|---------|
| 1 | Browsed http://aap.org | 31 |

A.2. Primary Study Search Strategies: Central Line-associated Bloodstream Infections (CLABSI) (May 5, 2021)

Table 3 Primary Search of MEDLINE: CLABSI

| # | Search History | Results |
|----|--|---------|
| 1 | exp Intensive Care Units, Neonatal/ or exp Intensive Care, Neonatal/ | 17500 |
| 2 | exp Infant, Newborn/ | 609494 |
| 3 | 1 or 2 | 610861 |
| 4 | exp Catheters, Indwelling/ | 19234 |
| 5 | exp Catheterization, Central Venous/ or exp Catheterization, Peripheral/ | 24828 |
| 6 | exp Umbilical Arteries/ or exp Umbilical Veins/ | 17948 |
| 7 | 4 and 6 | 157 |
| 8 | 5 and 6 | 303 |
| 9 | 4 or 5 | 39634 |
| 10 | 7 or 8 | 402 |
| 11 | PICC.mp. | 974 |
| 12 | Broviac.mp. | 364 |
| 13 | 9 or 10 or 11 or 12 | 40041 |
| 14 | exp Infection Control/ | 61617 |
| 15 | exp Cross Infection/ or exp Catheter-Related Infections/ | 60971 |
| 16 | exp Infusions, Intravenous/ae, mo [Adverse Effects, Mortality] | 1143 |
| 17 | exp Injections, Intravenous/ae, co, mo [Adverse Effects, Complications, Mortality] | 1300 |

Page 5 of 137

| 18 | 16 or 17 | 2409 |
|----|---|--------|
| 19 | 14 or 15 or 18 | 112730 |
| 20 | 3 and 13 and 19 | 425 |
| 21 | limit 20 to (English language and humans) | 385 |
| 22 | exp Bacteremia/ | 28376 |
| 23 | 19 or 22 | 137107 |
| 24 | 3 and 13 and 23 | 490 |
| 25 | limit 24 to (English language and humans) | 442 |
| 26 | 21 or 25 | 442 |
| 27 | limit 26 to yr="2012 -Current" | 150 |

Table 4 Primary Search of EMBASE: CLABSI

| # | Search History | Results |
|----|--|---------|
| 1 | Exp newborn intensive care/ or exp newborn/ | 385215 |
| 2 | Exp indwelling catheter/ or exp central venous catheter/ or exp catheterization/ | 162190 |
| 3 | Exp umbilical artery catheter/ or exp umbilical artery catheterization/ | 389 |
| 4 | Exp umbilical vein/ | 12348 |
| 5 | 2 and 4 | 342 |
| 6 | 2 or 3 or 5 | 162291 |
| 7 | Exp infection control/ or exp hospital infection/ or exp cross infection/ | 130845 |
| 8 | Exp bloodstream infection/ or exp catheter infection/ | 23173 |
| 9 | 7 or 8 | 149431 |
| 10 | 1 and 6 and 9 | 658 |
| 11 | Limit 10 to (English language and humans and embase) | 411 |

Table 5 Primary Search of Cochrane Library: CLABSI

| # | Search History | Results |
|---|--|---------|
| 1 | MeSH descriptor Intensive Care, Neonatal explode all trees | 120 |
| 2 | MeSH descriptor Intensive Care Units, Neonatal explode all trees | 84 |
| 3 | MeSH descriptor Infant, Newborn explode all trees | 153 |
| 4 | 1 or 2 or 3 | 206 |
| 5 | MeSH descriptor Catheters, Indwelling explode all trees | 46 |

| 6 | MeSH descriptor Catheterization, Central Venous explode all trees | 59 |
|----|---|----|
| 7 | MeSH descriptor Catheterization, Peripheral explode all trees | 52 |
| 8 | 5 or 6 or 7 | 91 |
| 9 | MeSH descriptor Umbilical Arteries explode all trees | 9 |
| 10 | MeSH descriptor Umbilical Veins explode all trees | 11 |
| 11 | 9 or 10 | 16 |
| 12 | 8 and 11 | 2 |
| 13 | 8 or 12 | 91 |
| 14 | 4 and 13 | 19 |

Table 6 Primary Search of CINAHL: CLABSI

| # | Search History | Results |
|----|--|---------|
| 1 | (MH "Infant, Newborn+") or (MH "Intensive Care, Neonatal+") or (MH "Intensive Care Units, Neonatal") | 78909 |
| 2 | MH "Central Venous Catheters+" | 2595 |
| 3 | (MH "Catheterization, Peripheral+") or (MH "Catheterization, Central Venous+") | 4398 |
| 4 | (MH "Umbilical Arteries") or (MH "Umbilical Veins") | 707 |
| 5 | 2 or 3 | 6420 |
| 6 | 4 and 5 | 39 |
| 7 | 5 or 6 | 6420 |
| 8 | MH "Infection Control+" | 46282 |
| 9 | (MH "Cross Infection+") or (MH "Catheter-Related Infections") | 23582 |
| 10 | MH "Bacteremia" | 3178 |
| 11 | (MH "Infusions, Intravenous/AE") or (MH "Infusions, Parenteral/AE") | 246 |
| 12 | 8 or 9 or 10 or 11 | 61658 |
| 13 | 1 and 7 and 12 | 215 |
| 14 | Limit 13 to (English language and human; exclude MEDLINE records) | 206 |

A.3. Primary Study Search Strategies: Central Line-associated Bloodstream Infections and Chlorhexidine (May 5, 2021)

Table 7 CLABSI and Chlorhexidine Search Strategy for MEDLINE

| # | Search History | Results |
|---|----------------|---------|

| 1 | exp Intensive Care Units, Neonatal/ or exp Intensive Care, Neonatal/ | 17500 |
|----|--|--------|
| 2 | exp Infant, Newborn/ | 609494 |
| 3 | 1 or 2 | 610861 |
| 4 | exp Catheters, Indwelling/ | 19234 |
| 5 | exp Catheterization, Central Venous/ or exp Catheterization, Peripheral/ | 24828 |
| 6 | PICC.mp. | 974 |
| 7 | Broviac.mp. | 364 |
| 8 | 4 or 5 or 6 or 7 | 40041 |
| 9 | exp Infection Control/ | 61617 |
| 10 | exp Cross Infection/ or exp Catheter-Related Infections/ | 60971 |
| 11 | exp Infusions, Intravenous/ae, mo [Adverse Effects, Mortality] | 1143 |
| 12 | exp Injections, Intravenous/ae, co, mo [Adverse Effects, Complications, Mortality] | 1300 |
| 13 | exp Bacteremia/ | 28376 |
| 14 | 9 or 10 or 11 or 12 or 13 | 137107 |
| 15 | Chlorhexidine.mp. or exp Chlorhexidine/ | 11575 |
| 16 | 3 and 15 | 326 |
| 17 | 15 and 8 and 14 | 290 |
| 18 | 16 or 17 | 590 |
| 19 | limit 18 to (English language and humans) | 535 |

Table 8 Primary Search of EMBASE: CLABSI and Chlorhexidine

| # | Search History | Results |
|---|--|---------|
| 1 | Exp newborn intensive care/ or exp newborn/ | 385215 |
| 2 | Exp indwelling catheter/ or exp central venous catheter/ or exp catheterization/ | 162190 |
| 3 | Exp umbilical artery catheter/ or exp umbilical artery catheterization/ | 389 |
| 4 | 2 or 3 | 162291 |
| 5 | Exp infection control/ or exp hospital infection/ or exp cross infection/ | 130845 |
| 6 | Exp bloodstream infection/ or exp catheter infection/ | 23173 |
| 7 | 5 or 6 | 149431 |
| 8 | 4 and 7 | 9679 |
| 9 | Exp chlorhexidine/ or chlorhexidine | 17183 |

| 10 | 1 and 9 | 420 |
|----|--|------|
| 11 | 8 and 9 | 852 |
| 12 | 10 or 11 | 1224 |
| 13 | Limit 12 to (English language and humans and embase) | 744 |

Table 9 Search of the Cochrane Library: CLABSI and Chlorhexidine

| # | Search History | Results |
|----|---|---------|
| 1 | MeSH descriptor Intensive Care, Neonatal explode all trees | 120 |
| 2 | MeSH descriptor Intensive Care Units, Neonatal explode all trees | 84 |
| 3 | MeSH descriptor Infant, Newborn explode all trees | 153 |
| 4 | 1 or 2 or 3 | 206 |
| 5 | MeSH descriptor Catheters, Indwelling explode all trees | 46 |
| 6 | MeSH descriptor Catheterization, Central Venous explode all trees | 59 |
| 7 | MeSH descriptor Catheterization, Peripheral explode all trees | 52 |
| 8 | 5 or 6 or 7 | 91 |
| 9 | MeSH descriptor Umbilical Arteries explode all trees | 9 |
| 10 | MeSH descriptor Umbilical Veins explode all trees | 11 |
| 11 | 9 or 10 | 16 |
| 12 | 8 and 11 | 2 |
| 13 | 8 or 12 | 91 |
| 14 | 4 and 13 | 19 |
| 15 | MeSH descriptor Chlorhexidine explode all trees | 88 |
| 16 | 13 and 15 | 11 |
| 17 | 4 and 15 | 8 |
| 18 | 16 or 17 | 12 |

B. Study Exclusion Criteria

Criteria for excluding studies from the literature review are:

1. Not relevant to key question

- 2. Not primary research
- 3. Meeting abstract only
- 4. No full text available
- 5. Not in English
- 6. No NICU patients included in study
- 7. Mixed patient population without NICU patient subgroups
- 8. Methods paper on HAI surveillance only
- 9. Descriptive epidemiology study only
- 10. Studies examining only non-modifiable risk factors for infection
- 11. Studies that do not provide a clear description of intervention and statistical analysis comparing time points before and after N<10 NICU patients with Outcome Definitions of interest (does not apply to studies evaluating severe adverse events such as death or permanent disfiguration)
- 12. Study only examining treatments of CLABSI
- 13. Study only examining catheter removal for documented CLABSIs
- 14. Study only examining peripheral IVs (note: this does not include Midline or PICCs)
- 15. Study with only endocarditis as a reported clinical outcome

Page **10** of **137**

C. Evidence Review

C.1. Non-sterile Gloves

Question 1. In NICU patients requiring a central line catheter, does the use of non-sterile gloves after hand hygiene compared with hand hygiene alone prior to every patient contact prevent CLABSI?

Table 10 The Summary of Evidence for Using Non-Sterile Gloves After Hand Hygiene vs. Hand Hygiene Alone Prior to Every Patient Contact to Prevent CLABSI

| Outcome | Findings | Quantity and Type of Evidence (Sample Size) | GRADE of Evidence for Outcome (Limitations of the Evidence) |
|-------------------|--|---|--|
| CLABSI* | • One single center RCT compared non-sterile glove use after hand hygiene with hand hygiene alone prior to every patient contact and found no difference in CLABSI rate (1.9 vs. 1.7, Rate Ratio: 0.90 (95% CI: 0.22-3.61), p = 0.88). | 1 RCT N=120 lines ¹ | Moderate • Imprecision: only one study |
| Possible CLABSI* | • One single center RCT compared non-sterile glove use after hand hygiene with hand hygiene alone prior to every patient contact and reported a decrease in possible CLABSI rate (9.4 vs. 3.4, Rate Ratio: 0.36 (95% CI: 0.16-0.81), p = 0.01). | 1 RCT N=120 lines ¹ | Moderate • Imprecision: only one study |
| BSI* | • One single center RCT compared non-sterile glove use after hand hygiene with hand hygiene alone prior to every patient contact and found no difference in BSI incidence (20/60 (33%) vs 14/60 (23%), difference in proportion: -10% (95% CI: -26 to 6), p = 0.22). | 1 RCT N=120 lines ¹ | Moderate • Imprecision: only one study |
| Gram Positive BSI | One single center RCT compared non-sterile glove use after hand hygiene with hand hygiene alone prior to every patient contact and reported a reduction in gram positive BSI incidence (19/60 [32%] vs. 9/60 [15%], Difference in proportion: -17% (95% CI: -31 to -1), p = 0.03). | 1 RCT N=120¹ lines¹ | Moderate • Imprecision: only one study |
| Gram Negative BSI | • One single center RCT compared non-sterile glove use after hand hygiene with hand hygiene alone prior to every patient contact and found no difference in gram negative BSI incidence (3/60 (5%) vs. 5/60 (8%), Difference in proportion: 3% (95% CI: -7 to 14), p = 0.46). | 1 RCT N=120¹ lines¹ | Moderate • Imprecision: only one study |

Table 11 Extracted Information for Non-Sterile Gloves After Hand Hygiene to Prevent CLABSI

| Study Information | Population and Setting | Intervention/ Study Groups | Definitions | Results |
|-------------------------------------|--------------------------------|---|--|--|
| Author: Kaufman ¹ | Number of Patients: N=120 | Intervention: n=60 | Outcome Definitions: | Primary Outcomes: |
| | Randomized N=124 | Group A: Glove use + HH | CLABSI: Centers for Disease Control and | CLABSI rate per 1000-line days: |
| Year: 2014 | Number of Lines: 120 lines | Non-sterile glove use after hand hygiene (HH) prior | Prevention definition (2008) | • Glove use + HH: 1.7 • HH Only: 1.9 |
| Study Design: Randomized control | Setting: NICU | to all contact with the patient, inside the bed | Possible CLABSI: detection of ≥1 blood cultures of any organism, and the | • Ratio: 0.90 (95% CI: 0.22-3.61) • p = 0.88 |
| trial | Location: US | area, and with all central and peripheral venous | presence of a central line within 72 hours in the absence of another source of | CLABSI, n/N (%): |
| Risk of Bias: Moderate | Dates: December 2008-June 2011 | catheters | infection | Glove use + HH: 4/60 (6.7%)HH only: 4/60 (6.7%) |

| **A-week minimum intervention duration after bedside of all birth; extended if infant required intravenous access (peripheral or central), or if line was removed and then subsequently needed infant admitted to the University NICU were eligible for the study if they had a birth weight All healthcare, professionals followed the Standard preventive measures: **All healthcare, professionals followed the Standard preventive measures: **All healthcare, used non-streing downs for contact with brought of hygiene in healthcare, used non-streing lowes for contact with body fluids, used sterile gloves for easiening lowes were used when accessing arterial lines. **CLASS blundle for placement, maintenance, and removal of catheters** **CLASS blundle for placement, maintenance, and removal of catheters** **CLASS blundle for placement, maintenance, and removal of catheters** **Fluornazoide prophylaxis for all infants who is considered with clinical signs, and symptoms of infection and treated with clinical signs, and symptoms of infection and treated with clinical signs, and symptoms of infection and treated with clinical signs, and symptoms of infection and treated with clinical signs, and symptoms of infection and treated with clinical signs, and symptoms of infection and treated with clinical signs, and symptoms of infection and treated with clinical signs, and symptoms of infection and treated with clinical signs, and symptoms of infection and treated with clinical signs, and symptoms of infection and treated with clinical signs, and symptoms of infection and treated with clinical signs, and/or necessary symptoms of light symptoms of fluid (CSF) incletions; growth the standard preventive measures: **All healthcare, professionals followed the symptoms of the standard preventive measures: **All healthcare, professionals followed the symptoms of the standard prevention: **Hornity: 20/60 (33%) **Clove use + HH: 19/60 (12%) | Study Information | Population and Setting | Intervention/ Study Groups | Definitions | Results |
|---|-------------------|-----------------------------|---|--|---|
| birth; extended if infant required intravenous access (peripheral or central), or if line was removed and then subsequently needed intervention: n=60 Group B: H norm or outborn (preterm) infants admitted to the University NICU were eligible for the study if they had a birth weight 4:1000 gor gestational age <29 weeks and were <6 days old Exclusion Criteria: All inhorn or outborn Criteria: All inhorn or outborn (preterm) infants admitted to the University NICU were eligible for the eligible (2006) gor gestational age <29 weeks and were <6 days old Exclusion Criteria: NR Device/agent: NA Monitoring intervention: Hand hygiene compliance standard preventive measures: - All healthcare professionals followed the 5 moments of hand hygiene from the World Health Organization guidelines for hand hygiene in healthcare, used non-sterile gloves for contact with body fluids, used sterile gloves for sapptic procedures - For both groups, non-sterile gloves were used when accessing arterial lines - CLASS bundle for polacement, maintenance, and removal of clarabeters in lines - CLASS bundle for polacement, maintenance, and removal of clarabeters - Fluctomazule prophysiaxis - For both groups, non-sterile gloves for contact with body fluids, used sterile gloves for polacement, maintenance, and removal of clarabeters - Fluctomazule prophysiaxis - Fluctomazule prophysia | | • 4-week minimum | Signs were placed on a | Symptomatic BSIs: growth in ≥1 blood | • p = NR |
| access (peripheral or central), or if line was removed and then subsequently needed inclusion Criteria: All inborn or outborn (preterm) infants admitted to the University NICU were eligible for the study if they had a birth weight Control/Comparison: Pre-intervention: n=60 (Silve use + HH: 3.4 and/or NEC associated with clinical signs and symptoms of infection and treated with antimicrobial sand with the total University NICU were eligible for the study if they had a birth weight Control/Comparison: Pre-intervention: n=60 (Silve use + HH: 3.4 and/or NEC associated with clinical signs and symptoms of infection and treated with antimicrobial sand with the total platient, bed, and/or catheter (all central and peripheral venues catheters) contact with antimicrobial sand ware Control/Comparison: Pre-intervention: His of Mystem (HH) alone prior to all platient, bed, and/or catheter (all central and peripheral venues times (LCS) infections: growth of bacteria or fungi from ≥ 1 cultures or fungi | | intervention duration after | stand at the bedside of all | culture and treated | |
| access (peripheral or central), or if line was removed and then subsequently needed inclusion Criteria: All inborn or outborn (preterm) infants admitted to the University NICU were eligible for the study if they had a birth weight 1.000 or greatrational age 2.000 or gestational age | | birth; extended if infant | enrolled patients (with a | | · |
| protocol. SSJ, urinary tract infection, menigitis, and symptoms of infection and treated with clinical signs, and symptoms of infection and treated with admirted to the University NICU were eligible for the study if they had a britth weight < 1000g or gestational age <29 weeks and were <8 days old Exclusion Criteria: NR Standard preventive measures: All healthcare professionals followed the 5 moments of hand hygiene compliance All healthcare professionals followed the 5 moments of hand hygiene from the World Health Organization guidelines for hand hygiene from the World Health Organization guidelines for hand hygiene for the the survey in the maccessing arterial lines CLASSI bundle for placement, maintenance, and removal of catheters | | required intravenous | box of gloves) indicating | Late-onset invasive infection: > 72 hours | • Glove use + HH: 3.4 |
| removed and then subsequently needed Inclusion Criteria: All inborn or outborn [preterm] infants admitted to the University NICU were eligible for the study if they had a birth weight 4-Inverter of the study if they had a birth weight 4-Inverter of the study if they had a birth weight 4-Inverter of the study if they had a birth weight 4-Inverter of the study if they had a birth weight 4-Inverter of the study if they had a birth weight 4-Inverter of the study if they had a birth weight 4-Inverter of the study if they had a birth weight 4-Inverter of the study if they had a birth weight 4-Inverter of the study if they had a birth weight 4-Inverter of the study if they had a birth weight 4-Inverter of the study if they had a birth weight 4-Inverter of the study if they had a birth weight 4-Inverter of the study if they had a birth weight 4-Inverter of the study if they had a birth weight 4-Inverter of the study if they had a birth weight 4-Inverter of the study if they had a birth weight 4-Inverter of the study if they had a birth weight 4-Inverter of the study if they had a birth weight 4-Inverter of the study if they had a birth weight 4-Inverter of the study if they had a birth weight 4-Inverter of the study if they had a birth weight 4-Inverter of the study if they had a birth weight 4-Inverter of the study if they had a birth weight 4-Inverter of the study if they had a birth weight 4-Inverter of the study if they had a birth weight 4-Inverter of the study if they had a birth weight 4-Inverter of the study if they had a birth weight 4-Inverter of the study if they had a birth weight 4-Inverter of the study if they had a birth weight 4-Inverter of the study | | 15 5 | | | • HH Only: 9.4 |
| Subsequently needed Inclusion Criteria: All inborn or outborn [preterm] infants admitted to the University NICU were eligible for the study if they had a birth weight < 1000g or gestational age <29 weeks and were <8 days old Exclusion Criteria: NR Monitoring intervention: Hand hygiene compliance Standard prevention: Hand hygiene compliance or professionals followed the 5 moments of hand hygiene for mith world Health Organization guidelines for hand hygiene for mate whorld Health Organization guidelines for hand hygiene for the study fifty flags, used sterile glows for contact with body fluids, used sterile glows for contact with body fluids, used sterile glows were used when accessing arterial lines **CLASIS blundle for placement, maintenance, and removal of Catheters** **Prethervention: Hand hygiene (HH) alone pripheral with antimicrobials soap to fluid (CSF) infections: growth of bacteria or fungif from 2 1 cultures **Contract with catheter: whenever there was central and peripheral venous line.** **Contract with catheter: whenever there was central and peripheral venous line.** **All healthcare professionals followed the 5 moments of hand hygiene for healthcare, used non-sterile glows for contact with body fluids, used sterile glows for contact with body fluids, used sterile glows for contact with body fluids, used sterile glows are used when accessing arterial lines **CLASIS blundle for placement, maintenance, and removal of Catheters* **Fluornazole prophylaxis** **CLASIS blundle for placement, maintenance, and removal of Catheters* **Fluornazole prophylaxis** **CLASIS blundle for placement, maintenance, and removal of Catheters* **Fluornazole prophylaxis** **Presence of NEC: stage II or greater.** **Sampling, /Testing strategy: Blood and urine cultures* **Olive use + HH: 17 **He only: 20/60 (33-3)* **HI Only: 20/60 (33-3)* **HI Only: 20/60 (33-3)* **HI Only: 20/60 (33-3)* **HI Only: 20/60 (33-3)* **Difference in proportion: -10% (95% CI: -26 to 6) **To 6) **Difference in proportion: | | ** | protocol. | , | • Ratio: 0.36 (95% CI: 0.16-0.81) |
| Inclusion Criteria: All inborn or outborn [preterm] infants admitted to the University NICU were eligible for the study if they had a birth weight 1 Honly: 20/60 (33.3%) Inclusion Criteria: All inborn or outborn [preterm] infants admitted to the University NICU were eligible for the study if they had a birth weight 1 Honly: 20/60 (33.3%) Inclusion Criteria: NR Blood (BSI), urine (UTI), cerebrospinal fluid (CSF) interions: growth of bacteria or fluid (CSF) interion: growth of bacteria or fluid (CSF) interion: growth of bacteria or surface and peripheral venous catheter cuntary and peripheral venous catheter or surface and peripheral venous catheter or | | | | | • p = 0.01 |
| Inclusion Criteria: All inborn or outborn [preterm] infants admitted to the University NICU were eligible for the study if they had a birth weight <1000g or gestational age <29 weeks and were <8 days old Exclusion Criteria: NR Monitoring intervention: Hand hygiene compliance Standard preventive measures: • All healthcare professionals followed the 5 moments of hand hygiene in healthcare, used non-sterile gloves for contact with body fluids, used sterile gloves for contact with body fluids, used sterile gloves for contact with body fluids, used sterile gloves for aseptic procedures • For both groups, non-sterile gloves for agentic professional are are professional are are professional are graphylaxis for the dealth of graphylaxis for the machine and they green from the whole accessing arterial lines • CLABSI bundle for placement, maintenance, and removal of catheters • Fluconazole prophylaxis few large and many professional prophylaxis few large and mon-sterile gloves were used when accessing arterial lines • CLABSI bundle for placement, maintenance, and removal of catheters • Fluconazole prophylaxis few large and mon-sterile gloves are used when accessing arterial lines • CLABSI bundle for placement, maintenance, and removal of catheters • Fluconazole prophylaxis few large and prove and the professional prophylaxis few large and prove a | | subsequently needed | · • | , . | |
| • Hand hygiene (HH) alone prior to all patient, bed, and/or catheter (all central and peripheral venous catheter contact with catheter is whenever the hand hygiene compliance. Standard preventive measures: • All healthcare professionals followed the 5 moments of hand hygiene in healthcare, used non-sterile gloves for contact with body fluids, used sterile gloves for paseptic procedures. • For both groups, nonsterile gloves for palacement, maintenance, and removal of catheters. • ClaBsi bundle for placement, maintenance, and removal of catheters. • ClaBsi bundle for placement, maintenance, and removal of catheters. • Fuluconazole prophylaxis • Hand hygiene (HH) alone prior to all patient, bed, and/or catheters of frong from ≥ 1 cultures • Central line (CL) days: days with umbilical central venous line, peripherally inserted central catheter, or surgical central venous line, peripherally inserted central catheter, or surgical central venous line, peripherally inserted central catheter, or surgical central venous line, peripherally inserted central catheter, or surgical central venous line, peripherally inserted central catheter, or surgical central venous line, peripherally inserted central catheter, or surgical central venous line, peripherally inserted central catheter, or surgical central venous line, peripherally inserted central central venous line, peripheral venous catheter contact and when making or breaking a connection with the bud when: (1) giving medications or flush, (2) changing tubing, (3) accessing an injection port, and hygiene from the World v | | | | with antimicrobials | 1 |
| admitted to the University NICU were eligible for the study if they had a birth weight < 1000g or gestational age < 29 weeks and were < 8 days old Exclusion Criteria: NR Device/agent: NA Monitoring intervention: Hand hygiene compliance Standard preventive measures: • All healthcare professionals followed the 5 moments of hand hygiene from the World Health Organization guidelines for hand hygiene in healthcare, used non-sterile gloves for contact with body fluids, used sterile gloves for contact with body fluids, used sterile gloves for contact with body fluids, used sterile gloves for aseptic procedures For both groups, non-sterile gloves for and removal of catheters Fluconazole prophylaxis Feculia (CSF) infections: growth of bacteria or funity from ≥ 1 cultures por funity from ≥ 1 cultures Contact with unit most principle and peripheral venous catheter (CIL) days: days with umbilical venous line, peripherally inserted central catheter; usence there or surgical central venous line catheter, or surgical central venous line catheter venous catheter contact and when making or breaking a connection with the bub when: (1) giving medications or flush, (2) changing tubing, (3) accessing an injection port, and (4) adding a device Hand hygiene: using alcohol hand rub or washing hands with antimicrobial soap (e.g., 2% chiokidine gluconate) For both groups, non-sterile gloves for aspetic procedures For both groups, non-sterile gloves for aspetic procedures For | | | , , | | • Glove use + HH: 8/60 (13.3%) |
| NICU were eligible for the study if they had a birth weight < 1000g or gestational age <29 weeks and were <8 days old Exclusion Criteria: NR Device/agent: NA Monitoring intervention: Hand hygiene compiliance measures: • All healthcare professionals followed the 5 moments of hand hygiene from the World Health Organization guidelines for hand hygiene in healthcare, used non-sterile gloves for contact with body fluids, used sterile gloves for aseptic procedures • For both groups, non-sterile gloves were used when accessing arterial lines • CLABSI bundle for placement, maintenance, and removal of catheters • Filuconazole prophylaxis fewell the study in the place of the | | | , , , | | • HH Only: 20/60 (33.3%) |
| study if they had a birth weight < 1000g or gestational age <2 9 weeks and were <8 days old Exclusion Criteria: NR Monitoring intervention: Hand hygiene compliance Standard preventive measures: • All healthcare professionals followed the 5 moments of hand hygiene from the World Health Organization guidelines for hand hygiene in healthcare, used non-sterile gloves for contact with body fluids, used sterile gloves for asterile gloves for asterile gloves were used when accessing arterial lines • CLABSI bundle for placement, maintenance, and removal or canbeters • Fulconazole prophylaxis for each steam of the placement, maintenance, and removal or canbeters • Fulconazole prophylaxis for each steam of the placement, maintenance, and removal or canbeters • Fulconazole prophylaxis for each steam of the placement, maintenance, and removal or canbeters • Fulconazole prophylaxis • Central line (CL) days: days with umbilical to Central line (CL) days: days with umbilicat catheter, or surgical central venous line. Central line (CL) days: days with umbilical to Central line (CL) days: days with umbilical to the venous line. Central line (CL) days: days with umbilical catheter, or surgical central venous line. Central line (CL) days: days with umbilical to the tentral venous line. Central line (CL) days: days with umbilical catheter, or surgical central venous line. Central line (CL) days: days with umbilical to the tentral venous line. Central line (CL) days: days with umbilical catheter, or surgical central venous line. Phenous catheters whenever there was central and peripheral venous line. Contact with tatheter: whenever there was central and peripheral venous line. Contact with tatheter: whenever there was central and and peripheral venous line. Phenous catheter contact and when making or breaking a connection with the hub when: (1) giving medications or flush, (2) changing tubing, (3) accessing an injection port, and (4) adding a device Hand hygiene: using alcohol hand rub or washing hands with antimicr | | | | _ · · · · · · · · · · · · · · · · · · · | • p = NR |
| <1000g or gestational age <29 weeks and were <8 days old were <8 days old Bexclusion Criteria: NR Exclusion Criteria: NR Monitoring intervention: Hand hygiene compliance Standard preventive measures: All healthcare professionals followed the 14 month bygiene from the World Health Organization guidelines for hand hygiene from the World Health Organization guidelines for hand hygiene in healthcare, used non-sterile gloves for contact with body fluids, used sterile gloves for setrile gloves for contact with body fluids, used sterile gloves for setrile gloves were used when accessing arterial lines CLABSI bundle for placement, maintenance, and removal or canbeters Fluconazole prophylaxis For both groups or placement, maintenance, and removal or canbeters Fluconazole prophylaxis For Difference in proportion: -10% (95% CI: -26 to 6) Contact with catheter: whenever there was central and peripherall venous line Contact with catheter: whenever there was central and peripheral venous catheter contact and when making or breaking and energy according to the washing medications or flush, (2) changing tubing, (3) accessing an injection port, and (4) adding a device Hand hygiene: using alcohol hand rub or washing hands with antimicrobial soap (e.g., 2% chlorhexidine gluconate) Fersence of NEC: stage II or greater. Sampling /Testing strategy: Blood and urine cultures Other notes: None Other notes: None SIST (gram-positive), n/N (%): • Glove use + HH: 14/60 (23%) • HIH only: 20/60 (33%) • Difference in proportion: -10% (95% CI: -26 to 6) p p = 0.22 BSI (gram-positive), n/N (%): • Glove use + HH: 14/60 (23%) • HH only: 19/60 (25%) • Difference in proportion: -17% (95% CI: -31 to -1) • Clove use + HH: 14/60 (23%) • HIH only: 30/60 (5%) • HIH | | o . | · · · | or fungi from ≥ 1 cultures | |
| weeks and were <8 days old Exclusion Criteria: NR Monitoring intervention: Hand hygiene compliance Standard preventive measures: **All healthcare professionals followed the 5 moments of hand hygiene from the World Health Organization guidelines for hand hygiene in healthcare, used non-sterile gloves for contact with body fluids, used sterile gloves for aseptic procedures ** For both groups, non-sterile gloves were used when accessing arterial lines **CLABSI bundle for placement, maintenance, and removal of catheters **CLABSI bundle for placement, maintenance, and removal of catheters **Leucanazole prophylaxis feastle lighted rathers **Leucanazole prophylaxis feastle lighted rathers in the contact and when making or breaking a connection with the hub when: **CLABSI bundle for placement, maintenance, and removal of catheters **Leucanazole prophylaxis feastle lighted rathers **Leucanazole prophylaxis feastle leucanazole prophylaxis feastle lighted rathe | | | | Control line (CI) down down with week line | |
| Device/agent: NA Monitoring intervention: Hand hygiene compliance Standard preventive measures: • All healthcare professionals followed the 5 moments of hand hygiene in healthcare, used non-sterile gloves for contact with body fluids, used sterile gloves for aseptic procedures • For both groups, non- sterile gloves were used when accessing arterial lines • CLABSI bundle for placement, maintenance, and removal of catheters • Fluconazole propphylaxis feedlaged and the contact with catheter: whenever there was central and peripheral venous catheter contact and when making or breaking a connection with the hub when: (1) giving medications or flush, (2) changing tubing, (3) accessing an injection port, and (4) giving medications or flush, (2) changing tubing, (3) accessing an injection port, and (4) pe = 0.03 Hand hygiene: using alcohol hand rub or washing hands with antimicrobial soap (e.g., 2% chlorhexidine gluconate) Presence of NEC: stage II or greater. Sampling /Testing strategy: Blood and urine cultures Other notes: None Other notes: None Altheath care, used mon-sterile gloves for aseptic procedures • For both groups, non- sterile gloves were used when accessing arterial lines • CLABSI bundle for placement, maintenance, and removal of catheters • Fluconazole propphylaxis feedlaged and with antimicrobial soap (e.g., 2% chlorhexidine gluconate) Presence of NEC: stage II or greater. Sampling /Testing strategy: Blood and urine cultures Other notes: None Other notes: None Other notes: None | | | venous catheters] contact | | • Glove use + HH: 14/60 (23%) |
| Exclusion Criteria: NR Monitoring intervention: Hand hygiene compliance Standard preventive measures: • All healthcare professionals followed the 5 moments of hand hygiene from the World Health Organization guidelines for hand hygiene in healthcare, used non-sterile gloves for contact with body fluids, used sterile gloves were used when accessing arterial lines • CLABSI bundle for placement, maintenance, and removal of catheters • Filuconazole prophylaxis fersell wife for a where is a contact and when waking or breaking a connection with the hub when: (1) giving medications or flush, (2) changing tubing, (2) changing tubing, (2) changing tubing, (3) accessing an injection port, and (4) adding a device Hand hygiene compliance was central and peripheral venous catheters: whenever there was central and peripheral venous catheters whenever there was central and peripheral venous catheters whenever there was central and peripheral venous catheters: whenever there was central and peripheral venous catheters: whenever there was central and peripheral venous catheter swhenever there was central and peripheral venous catheter swhenever there was central and peripheral venous catheter swhenever there was central and peripheral venous catheter contact and when making or breaking a connection with the hub when: (1) giving medications or flush, (2) changing tubing, (3) accessing an injection port, and (4) adding a device Hand hygiene: using alcohol hand rub or washing hands with antimicrobial soap (e.g., 2% chlorhexidine gluconate) • Difference in proportion: -10% (9)% CI: -20 to 6) • p = 0.22 BSI (gram-positive), n/N (%): • Glove use + HH: 9/60 (35%) • Difference in proportion: -10% (9)% CI: -31 to -1) • p = 0.03 BSI (gram-negative), n/N (%): • Glove use + HH: 9/60 (35%) • Glove use + HH: 9/60 (35%) • Glove use + HH: 9/60 (8%) • Glove use + HH: 9/60 (55%) • Glove use + HH: 17 • Hand hy | | weeks and were <8 days old | 5 | | • HH only: 20/60 (33%) |
| Monitoring intervention: Hand hygiene compliance Standard preventive measures: • All healthcare professionals followed the 5 moments of hand hygiene from the World Health Organization guidelines for hand hygiene in healthcare, used non-sterile gloves for contact with body fluids, used sterile gloves for contact with body fluids, used sterile gloves for contact with body fluids, used sterile gloves were used when accessing arterial lines • CLABSI bundle for placement, maintenance, and removal of catheters • Fluconazole prophylaxis for sult infenterior with the hub when: (1) giving medications or flush, (2) changing tubing, (3) accessing an injection port, and (4) adding a device Hand hygiene: using alcohol hand rub or washing hands with antimicrobial soap (e.g., 2% chlorhexidine gluconate) (e.g., 2% chlorhexidine gluconate) Presence of NEC: stage II or greater. Sampling /Testing strategy: Blood and urine cultures Other notes: None Contact with catheter: whenever there was central and peripheral venous catheter contact and when making or breaking a connection with the hub when: (1) giving medications or flush, (2) changing tubing, (3) accessing an injection port, and (4) adding a device Hand hygiene: using alcohol hand rub or washing hands with antimicrobial soap (e.g., 2% chlorhexidine gluconate) (e.g., 2% chlorhexidine gluconate) Presence of NEC: stage II or greater. Sampling /Testing strategy: Blood and urine cultures Other notes: None Other notes: None To other notes: None Contact with catheter: whenever there was central making or breaking a connection with the hub when: (1) giving medications or flush, (2) changing tubing, (3) accessing an injection port, and (4) adding a device Hand hygiene: using alcohol hand rub or washing hands with antimicrobial soap (e.g., 2% chlorhexidine gluconate) (e.g., 2% chlorhexidine gluconate) Presence of NEC: stage II or greater. Sampling /Testing strategy: Blood and urine cultures Other notes: None To other notes: None To other notes: None To othe | | Evaluaion Cuitoria: ND | Device/agent: NA | catheter, or surgical central vehous line | • Difference in proportion: -10% (95% CI: -26 |
| was central and peripheral venous catheter contact and when making or breaking a connection with the hub when: • All healthcare professionals followed the 5 moments of hand hygiene from the World Health Organization guidelines for hand hygiene in healthcare, used non-sterile gloves for contact with body fluids, used sterile gloves for aseptic procedures • For both groups, nonsterile gloves were used when accessing arterial lines • CLABSI bundle for placement, maintenance, and removal of catheters • Fluconazole prophylaxis for the Willight and the procession and removal of catheters • Fluconazole prophylaxis • p = 0.22 BSI (gram-positive), n/N (%): • Glove use + HH: 9/60 (15%) • HH only: 19/60 (32%) • Difference in proportion: -17% (95% CI: -31 to -1) • p = 0.03 BSI (gram-negative), n/N (%): • Glove use + HH: 5/60 (8%) • HH only: 3/60 (5%) • Difference in proportion: -17% (95% CI: -7 to washing hands with antimicrobial soap (e.g., 2% chlorhexidine gluconate) • Presence of NEC: stage II or greater. Sampling /Testing strategy: Blood and urine cultures Other notes: None | | Exclusion Criteria: NR | BA it i i - t t | Contact with cathotor: who nover there | to 6) |
| catheter contact and when making or breaking a connection with the hub when: All healthcare professionals followed the 5 moments of hand hygiene from the World Health Organization guidelines for hand hygiene in healthcare, used non-sterile gloves for contact with body fluids, used sterile gloves for aseptic procedures For both groups, nonsterile gloves were used when accessing arterial lines CLABSI bundle for placement, maintenance, and removal of catheters Fluconazole prophylaxis catheter contact and when making or breaking a connection with the hub when: (1) giving medications or flush, (2) changing tubing, (3) accessing an injection port, and (4) adding a device Hand hygiene: using alcohol hand rub or washing hands with antimicrobial soap (e.g., 2% chlorhexidine gluconate) Presence of NEC: stage II or greater. Sampling /Testing strategy: Blood and urine cultures Other notes: None BSI (gram-positive), n/N (%): Glove use + HH: 9/60 (32%) Pilference in proportion: -17% (95% CI: -31 to -1) p = 0.03 BSI (gram-negative), n/N (%): Glove use + HH: 5/60 (3%) HH only: 3/60 (5%) Difference in proportion: -17% (95% CI: -7 to 14) p = 0.04 BSI rate per 100 study days: Glove use + HH: 17 HH only: 23 Risk Ratio: 0.63% (95% CI: 0.34 to 1.18%) p = 0.15 Late on-set infection (any BSI, UTI, CSF, or NEC), n/N (%): | | | _ | | • p = 0.22 |
| Standard preventive measures: • All healthcare professionals followed the 5 moments of hand hygiene from the World Health Organization guidelines for hand hygiene in healthcare, used non-sterile gloves for contact with body fluids, used sterile gloves for aseptic procedures • For both groups, nonsterile gloves were used when accessing arterial lines • CLABSI bundle for placement, maintenance, and removal of catheters • Fluconazole prophylaxis fersell is ferse this feature for the world. Standard preventive measures: • All healthcare professionals followed the 5 moments of hand hygiene dications or flush, (2) changing tubing, (3) accessing an injection port, and (4) adding a device Hand hygiene: using alcohol hand rub or washing hands with antimicrobial soap (e.g., 2% chlorhexidine gluconate) Hand hygiene: using alcohol hand rub or washing hands with antimicrobial soap (e.g., 2% chlorhexidine gluconate) Presence of NEC: stage II or greater. Sampling /Testing strategy: Blood and urine cultures Other notes: None breaking a connection with the hub when: (1) giving medications or flush, (2) changing tubing, (3) accessing an injection port, and (4) adding a device Hand hygiene: using alcohol hand rub or washing hands with antimicrobial soap (e.g., 2% chlorhexidine gluconate) • Glove use + HH: 5/60 (8%) • HI only: 3/60 (5%) • Hil only: 3/60 (5%) • Glove use + HH: 9/60 (15%) • Difference in proportion: -17% (95% CI: -31 to -1) • p = 0.03 BSI (gram-negative), n/N (%): • Glove use + HH: 9/60 (15%) • Difference in proportion: -17% (95% CI: -31 to -1) • p = 0.03 BSI (gram-negative), n/N (%): • Glove use + HH: 9/60 (15%) • Glove use + HH: 9/60 | | | Hand hygiene compliance | | |
| measures: • All healthcare professionals followed the 5 moments of hand hygiene from the World Health Organization guidelines for hand hygiene in healthcare, used non-sterile gloves for contact with body fluids, used sterile gloves for aseptic procedures • For both groups, non-sterile gloves were used when accessing arterial lines • CLABSI bundle for placement, maintenance, and removal of catheters • Fluconazole prophylaxis for all the antipulation and removal of catheters • For late the the service of the contact with body fluids, used sterile gloves for aseptic procedures • For both groups, non-sterile gloves were used when accessing arterial lines • CLABSI bundle for placement, maintenance, and removal of catheters • Fluconazole prophylaxis for all the professionals followed the 5 moments of hand hygiene tubing, (2) changing tubing, (3) accessing an injection port, and (4) adding a device Hand hygiene: using alcohol hand rub or washing hands with antimicrobial soap (e.g., 2% chlorhexidine gluconate) Presence of NEC: stage II or greater. Sampling / Testing strategy: Blood and urine cultures Sampling / Testing strategy: Blood and urine cultures Other notes: None • Glove use + HH: 19/60 (13%) • Hi Honly: 19/60 (32%) • Difference in proportion: -17% (95% CI: -31 to -1) • p = 0.03 BSI (gram-negative), n/N (%): • Glove use + HH: 5/60 (8%) • HH only: 3/60 (5%) • Difference in proportion: -17% (95% CI: -7 to 14) • p = 0.46 BSI rate per 100 study days: • Glove use + HH: 17 • HH only: 23 • Risk Ratio: 0.63% (95% CI: 0.34 to 1.18%) • p = 0.15 Late on-set infection (any BSI, UTI, CSF, or NEC), n/N (%): | | | Standard proventive | | BSI (gram-positive), n/N (%): |
| • All healthcare professionals followed the 5 moments of hand hygiene from the World Health Organization guidelines for hand hygiene in healthcare, used non-sterile gloves for contact with body fluids, used sterile gloves for aseptic procedures • For both groups, nonsterile glows were used when accessing arterial lines • CLABSI bundle for placement, maintenance, and removal of catheters • Fluconazole prophylaxis (1) giving medications or flush, (2) changing tubing, (3) accessing an injection port, and (4) adding a device (1) giving medications or flush, (2) changing tubing, (3) accessing an injection port, and (4) adding a device (2) changing tubing, (2) daysing medications or flush, (2) changing tubing, (3) accessing an injection port, and (4) adding a device (4) adding a device (4) adding a device (4) adding a device (5) brifference in proportion: -17% (95% CI: -31 to -1) (5) p = 0.03 BSI (gram-negative), n/N (%): (6) Glove use + HH: 5/60 (8%) (7) brifference in proportion: -17% (95% CI: -31 to -1) (8) p = 0.03 BSI (gram-negative), n/N (%): (9) p = 0.04 (1) p = 0.0 | | | • | 1 | • Glove use + HH: 9/60 (15%) |
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| Health Organization guidelines for hand hygiene in healthcare, used non-sterile gloves for contact with body fluids, used sterile gloves for aseptic procedures • For both groups, nonsterile gloves were used when accessing arterial lines • CLABSI bundle for placement, maintenance, and removal of catheters • Fluconazole prophylaxis for such as the state of the state | | | | | • p = 0.03 |
| guidelines for hand hygiene in healthcare, used non-sterile gloves for contact with body fluids, used sterile gloves for aseptic procedures • For both groups, non-sterile gloves were used when accessing arterial lines • CLABSI bundle for placement, maintenance, and removal of catheters • Fluconazole prophylaxis for hand hygiene: using alcohol hand rub or washing hands with antimicrobial soap (e.g., 2% chlorhexidine gluconate) Hand hygiene: using alcohol hand rub or washing hands with antimicrobial soap (e.g., 2% chlorhexidine gluconate) Presence of NEC: stage II or greater. Sampling /Testing strategy: Blood and urine cultures Sampling /Testing strategy: Blood and urine cultures Other notes: None Hand hygiene: using alcohol hand rub or washing hands with antimicrobial soap (e.g., 2% chlorhexidine gluconate) • HH only: 3/60 (5%) • Difference in proportion: 3% (95% CI: -7 to 14) • p = 0.46 SBI rate per 100 study days: • Glove use + HH: 17 • HH only: 23 • Risk Ratio: 0.63% (95% CI: 0.34 to 1.18%) • p = 0.15 Late on-set infection (any BSI, UTI, CSF, or NEC), n/N (%): | | | , , , | () and might be seen | · · |
| washing hands with antimicrobial soap (e.g., 2% chlorhexidine gluconate) washing hands with antimicrobial soap (e.g., 2% chlorhexidine gluconate) presence of NEC: stage II or greater. sampling /Testing strategy: Blood and urine cultures Sampling /Testing strategy: Blood and urine cultures Other notes: None Glove use + HH: 5/60 (8%) HH only: 3/60 (5%) Difference in proportion: 3% (95% CI: -7 to 14) p = 0.46 BSI rate per 100 study days: Glove use + HH: 5/60 (8%) HH only: 3/60 (5%) Difference in proportion: 3% (95% CI: -7 to 14) p = 0.46 BSI rate per 100 study days: Glove use + HH: 5/60 (8%) HH only: 3/60 (5%) Difference in proportion: 3% (95% CI: -7 to 14) p = 0.46 BSI rate per 100 study days: HH only: 23 Risk Ratio: 0.63% (95% CI: 0.34 to 1.18%) p = 0.15 Late on-set infection (any BSI, UTI, CSF, or NEC), n/N (%): | | | | Hand hygiene: using alcohol hand rub or | BSI (gram-negative), n/N (%): |
| used non-sterile gloves for contact with body fluids, used sterile gloves for aseptic procedures • For both groups, nonsterile gloves were used when accessing arterial lines • CLABSI bundle for placement, maintenance, and removal of catheters • Fluconazole prophylaxis for all infects when | | | | , , , | • Glove use + HH: 5/60 (8%) |
| contact with body fluids, used sterile gloves for aseptic procedures • For both groups, nonsterile gloves were used when accessing arterial lines • CLABSI bundle for placement, maintenance, and removal of catheters • Fluconazole prophylaxis for all infects when a contact with body fluids, used sterile gloves for aseptic procedures • CLABSI bundle for placement, maintenance, and removal of catheters • Fluconazole prophylaxis for all infects when accessing arterial lines • Difference in proportion: 3% (95% CI: -7 to 14) • p = 0.46 • Difference in proportion: 3% (95% CI: -7 to 14) • p = 0.46 • Clabsi bundle for placement, maintenance, and removal of catheters • Fluconazole prophylaxis for all infects when accessing arterial lines • Clabsi bundle for placement, maintenance, and removal of catheters • Fluconazole prophylaxis for all infects when accessing arterial lines • Clabsi bundle for placement, maintenance, and removal of catheters • Fluconazole prophylaxis for all infects when accessing arterial lines • Difference in proportion: 3% (95% CI: -7 to 14) • p = 0.46 • BSI rate per 100 study days: • Glove use + HH: 17 • HH only: 23 • Risk Ratio: 0.63% (95% CI: 0.34 to 1.18%) • p = 0.15 Late on-set infection (any BSI, UTI, CSF, or NEC), n/N (%): | | | | | • HH only: 3/60 (5%) |
| used sterile gloves for aseptic procedures For both groups, nonsterile gloves were used when accessing arterial lines CLABSI bundle for placement, maintenance, and removal of catheters Fluconazole prophylaxis for all infents when | | | _ | | Difference in proportion: 3% (95% CI: -7 to |
| aseptic procedures • For both groups, nonsterile gloves were used when accessing arterial lines • CLABSI bundle for placement, maintenance, and removal of catheters • Fluconazole prophylaxis for all infeate when | | | | Presence of NEC: stage II or greater. | 14) |
| • For both groups, non- sterile gloves were used when accessing arterial lines • CLABSI bundle for placement, maintenance, and removal of catheters • Fluconazole prophylaxis for all infeate who | | | _ | | |
| when accessing arterial lines • CLABSI bundle for placement, maintenance, and removal of catheters • Fluconazole prophylaxis • Fluconazole prophylaxis • For all infeate whe | | | For both groups, non- | Sampling /Testing strategy: Blood and | , i |
| lines • CLABSI bundle for placement, maintenance, and removal of catheters • Fluconazole prophylaxis • or all infants who | | | | urine cultures | BSI rate per 100 study days: |
| • CLABSI bundle for placement, maintenance, and removal of catheters • Fluconazole prophylaxis • For all infants who | | | when accessing arterial | | • Glove use + HH: 17 |
| placement, maintenance, and removal of catheters • Fluconazole prophylaxis for all infants who | | | lines | Other notes: None | • HH only: 23 |
| placement, maintenance, and removal of catheters • Fluconazole prophylaxis for all infants who | | | CLABSI bundle for | | • Risk Ratio: 0.63% (95% CI: 0.34 to 1.18%) |
| and removal of catheters • Fluconazole prophylaxis for all infants who | | | placement, maintenance, | | |
| • Fluconazole prophylaxis for all infants who | | | and removal of catheters | | · · |
| for all infants who | | | Fluconazole prophylaxis | | |
| | | | for all infants who | | • Glove use + HH: 19/60 (32%) |
| weighed <1000g at birth | | | weighed <1000g at birth | | Page 12 of 137 |

| Study Information | Population and Setting | Intervention/ Study Groups | Definitions | Results |
|-------------------|------------------------|--|-------------|--|
| | | and/or had a gestational | | • HH only: 27/60 (45%) |
| | | age <28 weeks, or any | | • Difference in proportion: -12% (-28 to 6%) |
| | | infant with necrotizing | | • p = 0.13 |
| | | enterocolitis (NEC) or | | · · |
| | | gastroschisis | | Any infection rate per 100 study days: |
| | | Antibiotic stewardship | | • Glove use + HH: 27 |
| | | including limited use of | | • HH only: 35 |
| | | third- and fourth- | | • Risk Ratio: 0.67% (95% CI: 0.41 to 1.10%) |
| | | generation | | • p = 0.12 |
| | | cephalosporins and | | p 0.122 |
| | | carbapenems | | Topic-specific outcomes: |
| | | Limited use of postnatal | | Central line days / patient days (%): |
| | | corticosteroids, | | • Glove use + HH: 2,374/5,323 (44.6%) |
| | | histamineH2 receptor | | • HH only: 2,125/5,303 (40.1%) |
| | | blockers, and proton | | • p = 0.43 |
| | | pump inhibitors | | ν μ = 0.43 |
| | | Weekly changing of all | | Hand hygiene compliance, observed monthly |
| | | nasogastric and orogastric | | (%): |
| | | tubes | | • 2,675/3,385 (79%) |
| | | All patients with NEC were | | |
| | | placed in contact isolation | | Adverse events: NR |
| | | in which gowns and non- | | |
| | | sterile gloves were used | | |
| | | while patients were | | |
| | | receiving antimicrobials. | | |
| | | Auditing of compliance | | |
| | | performed throughout | | |
| | | the study | | |

Table 12 Risk of Bias of Randomized Controlled Trials on Using Non-Sterile Gloves After Hand Hygiene

| | Described as randomized | Randomization appropriately performed | Described as double- blind | Outcome assessor blinded | Study participant blinded | Investigator blinded | Attrition described | Attrition smaller than 10-15% of assigned patients | Attrition appropriately analyzed | Funding source(s) disclosed and no obvious conflict of interest | Overall Risk of Bias |
|------------------------------|-------------------------------|---|----------------------------------|--------------------------------|---------------------------------|-------------------------|------------------------|--|--|--|-------------------------|
| Kaufman 2014 ¹ | ✓ | √ | | | ~ | | ✓ | ✓ | | √ | Moderate |

Page 13 of 137

C.2. Central Line Type

Key Question 2: In NICU patients requiring central venous catheters, does the use of one central line catheter type, compared with another, prevent CLABSI?

Table 13 The Summary of Evidence on UVC vs. Peripheral Catheters to Prevent CLABSI

| Outcome | Findings | Quantity and Type of Evidence and Sample Size | GRADE of Evidence for Outcome and Limitations of the Evidence |
|--------------------------|---|--|---|
| CLABSI* | One observational study² reported a two-fold increase in the risk of CLABSI for UVCs compared with PICCs in a multivariable analysis (aHR 1.00 vs. 0.51 (95% CI: 0.40 – 0.66)). Two observational studies suggested no difference in the incidence of CLABSI when comparing UVC and PICCS. One observational study³ reported no difference in the incidence of catheter removal for CLABSI for UVCs compared to PICCs (15% vs 19%, p = NR). This result may have been confounded by shorter dwell time for UVCs compared with PICCs (6.9±2.7 vs 10.2±5.2, p <0.001). One observational study⁴ found no difference in the rate of CLABSI for UVC compared with PICCs (P = 0.952) | 3 OBS n= 3985 lines ² n=203 lines ³ n = 71 lines ⁴ | Very Low • Inconsistency: studies reporting different results |
| Catheter-associated BSI* | • One observational study reported no difference in the risk developing a CA-BSI per when comparing PICCs and UVCs (Adjusted IRR:1.18 (95% CI: 0.59–2.34); p = 64). | 1 OBS n=540 lines ⁵ | Very Low • Imprecision: only one study |
| Late Onset Sepsis* | • One observational study reported no difference the risk of developing a CA-BSI per when comparing PICCs and UVCs (Adjusted IRR: 1.06 (0.64–1.75); p = 82). | 1 OBS n=540 lines ⁵ | Very Low • Imprecision: only one study |
| Adverse Events | Two observational studies noted no difference in adverse events associated with both UVCs and PICCs including obstruction, extravasation, dislocation, and leakage. | 2 OBS n=203 lines ³ n = 71 lines ⁴ | Very Low ● Imprecision: only one study |

Table 14 The Summary of Evidence for the Efficacy of All Catheter Types to Prevent CLABSI

| Outcome | Findings | Quantity and Type of Evidence and Sample Size | GRADE of Evidence for Outcome and Limitations of the Evidence |
|---------|---|---|---|
| CLABSI* | One observational study⁶ found a higher incidence of CLABSI for tunneled catheters, PICC, and CVCs when compared with UVCs (p = 0.001); however in multivariable analysis, central line insertion in the operating theater (including CVCs and tunneled catheters) was a significant risk factor for CLABSI (OR 8.1 (95% CI 1.2 – 54.7); p = 0.03. One large multicenter observational study⁷ found the incidence of CLABSI for tunneled catheters was 2.4 times as high as the CLABSI incidence for PICCs (p<0.001). The accompanying median dwell time was shorter for PICCS than it was for tunneled catheters. One observational study⁸ reported a higher rate of CLABSI for PICCs than for extended dwell peripheral intravenous catheters (EPIV) (0 vs. 0.68/ 1000 days; p = NR) One observational study⁹ found no difference in the incidence of UAC, UVC, short duration venous catheter, PICC, and tunneled catheters (P = 0.816). | | Very Low • Inconsistency: studies reported different results |

| Outcome | Findings | Quantity and Type of Evidence and Sample Size | GRADE of Evidence for Outcome and Limitations of the Evidence |
|------------------------------|---|---|--|
| Catheter associated- BSI* | One observational study (de Brito 2010) reported a higher rate of catheter associated BSI for PICCs than for other catheters (including UVC, intracaths, and phlebotomy catheters) (p<0.01). | 1 OBS n = 461 ¹⁰ | Very Low ■ Imprecision: only one study, wide confidence intervals |
| Nosocomial BSI* | One observational study reported higher infection rates associated with percutaneous venous and tunneled catheters compared with UVCs (Crude RR: 1, p<0.05). | 1 OBS n=19,507 infants ¹¹ | Very Low • Imprecision: only one study |
| Nosocomial Sepsis* | One observational study reported higher sepsis incidence associated with tunneled and percutaneous catheters compared with umbilical catheters (p<0.0001). | 1 OBS n=3,107 lines ¹² | Very Low • Imprecision: only one study |
| Infiltration | One observational study found higher rates of infiltration associated with PICCs compared with UAC, UVC, short duration venous catheter, and tunneled catheters (IR: 12.4 CLABSI/ 1000 days). | 1 OBS n = 400 lines ⁹ | Very Low ● Imprecision: only one study |
| Adverse events | • One observational study reported a higher rate of obstruction, peritonitis, and premature ventricular contractions in infants with PICCs compared with EPIVs, however infants with EPIVs received a higher incidence of hyaluronidase treated IV fluid extravasation. | 1 OBS n = 2,828 patients ⁸ | Very Low ● Imprecision: only one study |

Table 15 Extracted Information on Central Line Type

| Study Information | Population and Setting | Intervention/ Study Groups | Definitions | Results |
|----------------------------------|---------------------------------|---|--|---------------------------------------|
| Author: | Number of patients: | Study Groups: | Outcome Definitions: | Primary Outcomes: |
| Konstantinidi ⁴ | N = 71 VLBW | Group A: n= 34 PICC (Because | CLABSI: CDC definition: Presence | CLABSI Rate/ 1000 line days: |
| | Number of lines: N=71 | UVC insertion failed during | of bacteria in a single blood culture (for | • PICC: 2.28 |
| Year: 2019 | | first 3 days of life) | organism not commonly present on | • UVC: 2.59 |
| | Setting: Tertiary NICU | Insertion was performed | the skin), or in two or more blood | • p = 0.952 |
| Study Design: | | during the morning shift | cultures (for organisms commonly | CLABSI Incidence: |
| Cohort study | Location: Greece | by a trained group of | present on the skin), obtained from a | • PICC: 1/34 (2.9%) |
| | | neonatologists and | symptomatic infant either within 48 h | • UVC: 1/37 (2.7%) |
| Risk of Bias: Moderate | Dates: 18 months (NR when) | nurses. The same group | after a central catheter insertion or within a 48-h period following | • p = 0.952 |
| | Inclusion Criteria: (1) Birth | was also responsible for | catheter removal, and not related to | Topic-specific outcomes: |
| | weight below 1500 g and | infant monitoring and | an infection at another site | Catheter dwell time mean±SD (days) |
| | gestational age < 32 weeks. | catheter removal. | | • PICC: 11.91 ± 6.93 |
| | Gestational age was defined by | Group B: n= 37 UVC only, no | Probable but unproven sepsis: Either | • UVC: 10.43±5.38 |
| | strict criteria, prioritizing | PICC insertion | clinical signs (aggravated clinical status | • p = 0.152 |
| | menstrual dating confirmed by | UVC access (with single- | presenting with apnea, hyperthermia | p 5.252 |
| | early ultrasound. (2) Insertion | lumen umbilical catheters) of | or hypothermia, tachycardia or | Adverse events: NR |
| | of CVC (UVC or PICC) in our | The inferior vena cava was | bradycardia, hypotension, | Obstruction, n/N (%) |
| | NICU. | performed by a group of | hyperglycemia), and/or on laboratory | • PICC: 1/34 (2.9%) |
| | | trained neonatologists | atologists Tindings (elevated C-reactive protein | • UVC: 0 |
| | Exclusion Criteria: | within the incubator, | along with two of the following: | Local edema +skin irritation, n/N (%) |
| | (1) Catheter removal within 24 | under sterile conditions. | Immature/mature white blood cell | • PICC: 2/34 (5.88 %) |
| | h following insertion because | ander sterne conditions. | ratio > 0.2, low (<100,000) platelet | • UVC: 0 |
| | of inappropriate line tip | | count, neutrophils white blood cell | |

| Study Information | Population and Setting | Intervention/ Study Groups | Definitions | Results |
|-------------------|--|--|--|--------------------------|
| | position, as the complication | Device/agent: Catheter type | count of <1500 without positive blood | Skin irritation, n/N (%) |
| | rate was expected to be low | | culture, and being defined as a | • PICC: 1/34 (2.9 %) |
| | due to the short indwelling | Standard preventive | systemic condition resulting from an | • UVC: 0 |
| | time; (2) CVC insertion in | measures: | adverse reaction to the presence of an | |
| | another center, because of | Choice of catheter was | infectious agent that was neither | |
| | possible differences or | based on protocol. | present nor incubating at the time of | |
| | incomplete data regarding the insertion procedure that might | • In VLBWs infants | admission to the hospital | |
| | affect the complication rate; | scheduled for a long NICU | Sampling /Testing strategy: | |
| | (3) congenital abnormality; | hospitalization, the | Whenever a neonate presented with | |
| | and (4) necrotizing | preferred option was | clinical signs or symptoms of sepsis, | |
| | enterocolitis (NEC) Bell stage II | catheter insertion in the | blood culture was performed prior to | |
| | or III, during the first five days | umbilical vein on the first | antibiotic therapy initiation. Blood | |
| | of life. | or second day of life. In | specimens were collected through | |
| | | case the first UVC | peripheral venipuncture, on separate | |
| | | insertion attempt in the | occasions: from at least two separate | |
| | | inferior vena cava failed | blood draws on the same or | |
| | | or in case of early UVC | consecutive calendar days, or two | |
| | | catheter removal due to | separate site preparations | |
| | | various reasons, a PICC | (decontamination steps) performed | |
| | | insertion was performed, | during specimen collection. No blood | |
| | | usually after the third day | specimens were drawn through | |
| | | of life. | Other notes: None | |
| | | Skin antiseptic | other notes. None | |
| | | preparation included | | |
| | | cleansing the site three | | |
| | | times with a cotton swab | | |
| | | remoistened with | | |
| | | povidone-iodine 10%. To | | |
| | | avoid prolonged exposure | | |
| | | to iodine, skin sites | | |
| | | disinfected with | | |
| | | povidone-iodine were | | |
| | | wiped with sterile normal | | |
| | | saline solution after 60 s | | |
| | | until all antiseptic stains | | |
| | | were removed. | | |
| | | The distal edge of the | | |
| | | catheter was disinfected | | |
| | | with a 0.5% | | |
| | | chlorhexidine/alcohol 70% | | |
| | | solution at least three | | |

| Study Information | Population and Setting | Intervention/ Study Groups | Definitions | Results |
|------------------------|----------------------------------|---|----------------------------------|--|
| | | times daily, according to | | |
| | | the instructions of the | | |
| | | Infectious Diseases | | |
| | | Committee of Hospital | | |
| Author: | Number of patients: | Study Groups: | Outcome Definitions: | Primary Outcomes: |
| Chenoweth ⁸ | N = 2,828 | All PIV: 2,828 | CLABSI: NR | CLABSI rate/ 1,000 line days |
| | Number of lines: N= NR | EPIV: n=432 | Complications: NR | • EPIV: 0 |
| Year: 2018 | | Neonates who are 32 | | • PICC: 0.68 |
| | Setting: Level III NICU | weeks of gestation or | Sampling /Testing strategy: None | • p = NA |
| Study Design: | | more and weighing 1500g | | |
| Prospective cohort | Location: USA | or more at birth with | Other notes: None | Topic-specific outcomes: |
| study | D | difficult or limited venous | | Catheter dwell time, mean (SD), days |
| B: 1 (B: | Dates: August 2012 – | access that is likely to be | | • EPIV 4.0 (2.3) |
| Risk of Bias: | December 2016 | required up to 4 weeks. | | • PICC: 7.31 (4.4) |
| Moderate | Inclusion Criteria: All neonates | Excluded: Neonates | | • p < 0.001 |
| | who were 32 weeks of | requiring fluid greater | | |
| | gestation or older and weighed | than dextrose 12.5% | | Adverse events: |
| | 1500 g or more at birth with | concentration, total | | Incidence of hyaluronidase treated IV fluid |
| | EPIV catheter, PICC, and/or PIV | parenteral nutrition | | extravasation, % |
| | catheter placements. | osmolarity greater than | | • EPIV: 1.2 |
| | catheter placements. | 900 mOsm/L, and/or | | • PIV: 3.9 |
| | Exclusion Criteria: NR | medications that are | | • p = 0.004 |
| | | | | |
| | | administrated via central | | Premature ventricular contractions, rate/ 1000 |
| | | catheters. | | catheter days |
| | | PICC: n=202 | | • EPIV: 0 |
| | | PICC Group inclusion | | • PICC: 0.68 |
| | | criteria: NR | | • p = NA |
| | | Device/agent: Catheter type | | Superior vena cava obstruction, rate/ 1000 |
| | | Device/agent: catheter type | | catheter days |
| | | Standard preventive | | • EPIV: 0 |
| | | measures: | | • PICC: 0.68 |
| | | Implemented a CLABSI | | • p = NA |
| | | | | Peritonitis rate/ 1000 catheter days |
| | | | | • EPIV: 0 |
| | | | | • PICC: 0.68 |
| | | | | • P = NA |
| | | | | Success rate (%) |
| | | | | • EPIV: 71.1 |

| Study Information | Population and Setting | Intervention/ Study Groups | Definitions | Results |
|-------------------------|--|--|---|--|
| | | | | • PICC: 83.6 |
| | | | | • p = 0.001 |
| Author: | Number of patients: | Case: | Outcome Definitions: | Primary Outcomes: |
| Geldenhuys ⁶ | N = 95 | CLABSI n=19 | HAI: CDC/NHSN 2014 definition used | CLABSI Rate (overall): |
| | Number of lines: N=95 | | | • 5.9/1 000 line days |
| Year: 2017 | | Control: | CLABSI: | CLABSI Incidence: |
| | Cases were significantly | Non-CLABSI n=76 | Laboratory-confirmed bloodstream | • UVC: 6/55 (10.9%) |
| Study Design: | younger in GA than control, | • 4 random controls were | infection (LC-BSI) in a patient with a | • PICC: 6/23 (26%) |
| Retrospective case | and had longer lengths of stay | selected for each case | central line in situ for at least 2 | • CVC: 4/14 (28%) |
| control study | and high lands | | calendar days (where line insertion is | • Tunneled: 3/3 (100%) (3 tunneled lines |
| Diel of Dies. | Setting: NICU and NICU wards | Device/agent: Catheter type | day 1). | inserted in 2-year period and all 3 |
| Risk of Bias: Low | Location: South Africa | Chandand annual the | LC-BSI occurred within 1 day of line | developed CLABSI) |
| LOW | Location: South Africa | Standard preventive measures: | removal | • p = 0.001 |
| | Dates: August 9, 2012 – July | | The definitions for HAI and LC-BSI | |
| | 31, 2014 | Implemented a CLABSI surveillance program, and | must be met before the definition of CLABSI can be applied, and other HAI | CLABSI Incidence by insertion setting: |
| | 31, 2014 | insertion and | must be excluded. | • NICU: 12/82 (14.6%) |
| | Inclusion Criteria: | maintenance bundles at | must be excluded. | • Theatre: 6/8 (75%) |
| | All cases within the 2-year | start of study (no baseline | CLABSI rate per 1000 central line days is | • Neonatal Ward: 1/5: (20%) |
| | study period | data) | calculated by dividing the number of | • p = 0.001 |
| | 4 randomly selected | UVCs and PICCs are | CLABSIs by the number of central line | • OR: 8.1 (95% CI 1.2 – 54.7) |
| | controls per CLABSI event | inserted by pediatric | days and multiplying the result by | • p = 0.03 |
| | were included. | registrars or medical | 1000. | Tonio anosifio autormos: |
| | Central line insertion | officers | CLABSI bundle: strategy for insertion and | Topic-specific outcomes: Catheter dwell time in NICU (incidence) Overall |
| | requirements include: | CVCs and Tunneled lines | maintenance of central lines, which | p = 0.007 |
| | Neonates who need | are inserted in patients in | includes several evidence-based best | < 4 days |
| | TPN and/or inotropes | whom intravenous access | practices implemented | • Case: 2/19 (11%) |
| | neonates who require | is difficult, where | simultaneously | • Control: 34/76 (45%) |
| | intravenous fluids | attempts at insertion of | Line days: total number of days of | 4 - 8 days |
| | and/or antibiotics | other central lines have | exposure to central venous catheters | • Case: 9/19 (47%) |
| | where peripheral | failed, and/or in post- | by all patients in the selected | • Control: 30/76 (39%) |
| | intravenous access is | surgical patients who | population and time period | > 8 days |
| | not possible or difficult | need TPN. | Adverse events: NA | • Case: 8/19 (42%) |
| | to obtain | Tunneled lines are | Sampling /Testing strategy: Blood | • Control: 12/76 (16%) |
| | Evolucion Critoria | inserted by the pediatric | cultures | |
| | Exclusion Criteria: Umbilical arterial lines | surgical team and CVCs by either the pediatric | | Time to CLABSI after line insertion (median IQR) |
| | Official afterial lifes | surgery or anesthetic | Other notes: | • UVC: 2 days (2-4) |
| | | team. | • Gram-negative pathogens were (54%) | • PICC: 9 days (6-13) |
| | | | dominant pathogens and half the | • CVC: 7 days (6-10) |
| | | | premature infants had surgery (stoma | • Tunneled: 20 (19-35) |
| | | | repairs) | |
| | | | | Catheter dwell time in NICU for CLABSI, |
| | | | | (median IQR) |

| Study Information | Population and Setting | Intervention/ Study Groups | Definitions | Results |
|--------------------------------|---|------------------------------------|---|---|
| | | | | All line types: 8 days (14-18) |
| | | | | • UVC: 4 days (3-5) |
| | | | | • PICC: 13 days (8-13) |
| | | | | • CVC: 8 days (8-11) |
| | | | | • Tunneled: 22 days (21-36) |
| | | | | Adverse events: NR |
| | | | | Attributable Mortality: |
| | | | | • 3/5 (60%) |
| Author: Sanderson ² | Number of patients: | Study Groups: | Outcome Definitions: | Primary Outcome: |
| | UVC only: 1392 | UVC only | CLABSI: | CLABSI Multivariable hazard ratio, aHR (95% CI) |
| Year: 2017 | PICC only: 1317 | (n=2668) | • (CDC, 2016) late onset sepsis (LOS) | • UVCs:1.00 |
| | • UVC & PICCs: 1276 | | with positive blood culture taken | • PICCs: 0.51 (0.40 – 0.66) |
| Study Design: | Number of Lines: | PICCs only | after the first 48 h of a CVC being in | • p = NR |
| Retrospective cohort | • UVC only: 1392 | (n = 3332) | situ | |
| study | PICC only: 1317 | | • (NSW Health criteria, 2008) 48 h of | CLABSI rate per 1000 days |
| | • UVC & PICCs: 1276 | Device/agent: Catheter type | CVC removal | UVCs: 9.88 CLABSI / 1000 days |
| Risk of Bias: | | | CLABSI episodes were assigned to the | PICCs: 9.09 CLABSI/ 1000 days |
| Low | Setting: Tertiary NICUs (n =10) | Standard preventive | CVC in situ according to this 48 h | • p = NR |
| | | measures: NR | post-insertion or post-removal cut- | |
| | Location: Australia | | off criteria if there were overlaps of | CLABSI incidence (% of catheter) |
| | | | CVC. | • UVCs: 116/ 2668 (4.3%) |
| | Dates: January 1, 2007 – | | | • PICCs: 287/ 3332 (8.6%) |
| | December 31, 2009 | | Incidence of CLABSI: number of episodes | • p < 0.01 |
| | | | / 1000 catheter-days and number of | |
| | Inclusion Criteria: | | episodes / 1000 catheters inserted. | Topic-specific outcomes: |
| | All infants: | | | Catheter days to CLABSI median, (IQR) |
| | Born within study period | | Early onset sepsis (EOS): positive blood | UVCs: 5.3 days (3.6, 7.3) |
| | Admitted to one of 10 | | culture in an infant taken within the first | PICCs: 8.1 days (5.2, 12.5) |
| | NICUs | | 48 hours of life and a clinical picture | • p < 0.01 |
| | with UVC or PICC inserted | | consistent with sepsis. | |
| | with 1st CVC insertion for ≥ | | Late enset sensis (LCC): | Adverse events |
| | 4 h | | Late onset sepsis (LOS): positive blood culture, clinical | NA |
| | • 1 or more CVCs inserted | | symptoms, and signs of sepsis and | |
| | throughout admission | | clinician decision to treat with antibiotics | |
| | during study period | | for ≥ 5 days, including coagulase- | |
| | | | negative staphylococci (CoNS) in the | |
| | Exclusion Criteria: | | Australian neonatal population, | |
| | CLABSIs occurring within | | (consistent with the definitions used by | |
| | the first 48 hours of life | 1 | (33 | |

| | İ | | | |
|-------------------------------|--|---|--|---|
| | , | | Neonatal Network and the Canadian | |
| | ļ | | Neonatal Network) | |
| | ļ | | | |
| | | | Causative pathogen: organism cultured | |
| | | | in the first episode of CLABSI of any CVC | |
| | | | | |
| | | | Adverse events: | |
| | | | NA | |
| | ļ | | Committee /Testing streets on a Discol | |
| | | | Sampling /Testing strategy: Blood | |
| | | | cultures | |
| | ļ | | Other notes: | |
| | | | Time to first CLABSI episode was used | |
| | | | if there were multiple CLABSI | |
| | | | episodes in the same CVC. The | |
| | | | primary outcome was the first CLABSI | |
| | | | in a UVC or PICC. | |
| | | | | |
| Author: Soares ⁹ N | Number of patients: | Study Groups: | Outcome Definitions: | Primary Outcomes: |
| N | N = 240 | Patients with infectious | Infectious complications: CLABSI: (CDC | CLABSI Rate (overall): 12.4 CLABSI/ 1000 days |
| Year: 2017 N | Number of lines: | central line complications n= | 2008 NHSN criteria) a primary | CLABSI Incidence (Overall): 48/240 (20%) |
| N | N= 400 central lines | 51 | bloodstream infection in a patient with a | |
| Study Design: | | | central line at the time or within 48-h | Infectious complications |
| - | Setting: Level III NICU, in a | Patients without infectious | period before the onset of sepsis clinical | • UACs: 3/55 (5.5%) |
| study re | regional hospital | central line complications n= | signs, without another identifiable | • UVC: 6/84 (7.1%) |
| | | 189 | infection source and with a positive | • Tunneled: 3/22 (13.6%) |
| Risk of Bias: Low Lo | Location: Portugal | | blood culture, collected when possible | • SDVC: 9/57 (15.8%) |
| | | Standard preventive | from central line. | • PICC: 30/182 (16.5%) |
| | Dates: July 1, 2014 – June 31, | measures: | | • p = 0.816 |
| 2 | 2016 | Radiograph obtained after | Line days to infection: number of days | |
| | In alveion Critonia | the last repositioning for | from line placement to onset of sepsis | Topic-specific outcomes: |
| | Inclusion Criteria: | CTP evaluation | signs | Length of catheter stay, (min-max) |
| | Admitted to NICU during study period who had a | Central lines were | CLABSI mortality: considered if cases | • UACs: 6 (2-28) |
| | study period who had a central line placed | removed due to elective | whose autopsy report referred to it | • UVC: 5 (2-18) |
| | central line placed | (end of therapy, discharge or death) or non-elective | whose autopsy report referred to it | • Tunneled: 16 (4-94) |
| _ | Exclusion Criteria: | reasons | Central venous catheters (UVC, PICC, | • SDVC: 11 (2-37) |
| | Neonates in NICU for less | Catheter removal because | Tunneled, and short duration venous | • PICC: 10 (2-46) |
| | than 3 days | of CLABSI is only required | catheter (SDVC)): central if the tip was | • p < 0.001 |
| | Neonates with central lines | if clinical deterioration | located at superior vena cava (SVC), | |
| | inserted and removed | after starting | inferior vena cava (IVC), or at SVC/IVC- | Adverse events |
| l l | same day | antibiotherapy or | right atrium junction and non-central if | Mortality rate: |
| | Same day | | | CLABSI related: 21.4% |

| Study Information | Population and Setting | Intervention/ Study Groups | Definitions | Results |
|-------------------|------------------------|---|--|--|
| | | persisting or relapsing bacteremia. • Tip culture follows central line removal | Length of catheter stay: the number of days the line stayed in the patient Central line utilization ratio: the number of catheter-days divided by the number of patient-days. Adverse events: Mechanical complications: occlusion, breakage, external leaking, infiltration, vasospasm, bleeding, phlebitis, exteriorization, pneumothorax, pericardial and pleural effusion, and cardiac tamponade Catheter related thromboembolism: catheter occlusion due to the presence of a thrombus; confirmed by echocardiography or ultrasonography. Occlusion: inability to infuse through a line or inability to flush it External leaking: a collection of intravenous fluid under the catheter dressing Infiltration: fluid extravasation into soft tissues and diagnosed by the inability to infuse fluid associated with swelling in the region of the catheter tip Phlebitis: inflammation tracking along the path of a non-occluded venous catheter expressed as tenderness, erythema, and/or induration at the surrounding area of the insertion site. Exteriorization: migration of the catheter until its tip surfaces Pleural or pericardial effusion: the escape of fluid from blood vessels and its collection, respectively, in pleural or pericardial space Sampling /Testing strategy: Blood cultures | Type of complications Mechanical UACs: 5/55 (9.1%) UVC: 6/84 (7.1%) Tunneled: 7/22 (31.8%) SDVC: 9/57 (15.8%) PICC: 45/182 (24.7%) p = 0.816 Infiltration UACs: 0/55 (0%) UVC: 0/84 (0%) Tunneled: 2/22 (9.1%) SDVC: 1/57 (1.8%) PICC: 28/182 (15.4%) p = 0.003 Rate of non-elective removals UACs: 7/55 (13.0%) UVC: 9/84 (11.7%) Tunneled: 7/22 (46.7%) SDVC: 11/57 (19.6%) PICC: 62/182 (39.5%) p < 0.001 |

Page **21** of **137**

| Author: Greenberg? N = 13,327 Year: 2015 Study Design: Retrospective cohort study Bisk of Bias: Low Dates: September 2011 – August 2013 August 2013 Infant with PICCs or tunneled catheters obtained from NCLABSI database during study dates I finant with PICCs or tunneled datheters obtained from NCLABSI database during study dates Peckusion Criteria: • Central lines inserted and removed within the first 2 days • Positive blood cultures or central lines when in freed the son current on the sittes • Positive blood culture was included in the analysis, and removed within the first 2 days • Positive blood cultures or recognized pathogens), only the first positive blood culture was included in the analysis, and removed within the first 2 days • Positive blood cultures or recognized pathogens), only the first positive blood culture was included in the analysis. • Central lines inserted and removed within the first 2 days • Positive blood cultures or central lines when infants achieved 120 m/L/kg per day of enteral feedings • Positive blood cultures or recognized pathogens, only the first positive blood culture was included in the analysis. • Central lines inserted and removed within the first 2 days • Positive blood cultures or central lines when infants achieved 120 m/L/kg per day of enteral feedings • Positive blood cultures or central lines when infants achieved 120 m/L/kg per day of enteral feedings • Positive blood cultures or recognized pathogen not related to an infection and another site or cannot a nonther site or infection and another site or site of single catheter type. Standard preventive measures: • Low Device/agent: Catheter type. Standard preventive measures: • Low Device/agent: Catheter type. Standard preventive measures: • Laks I sate lead to catheters: 3/9/1116 (3.5%) • PiCCS: 3/1/451 (1.4%) • PiCCS: 3/1/451 (1.5%) • PiCCS: 3/1/451 (1.5%) • PiCCS: 3/1/451 (1.6%) • PiCCS: |
|---|
| Other notes: None (0.8 − 12.0) • PICCs: 7/765 (0.9%); HR 1.5 (0.7− 3.2) Week 7 |

| Study Information | Population and Setting | Intervention/ Study Groups | Definitions | Results |
|-------------------|------------------------|----------------------------|-------------|--|
| | | | | • PICCs: 4/453 (0.9%); HR 1.4 (0.5-4.0) |
| | | | | Week 8 |
| | | | | • Tunneled catheters: 1/288 (0.4%); HR 1.3 (0.1-20.3) |
| | | | | • PICCs: 2/183 (1.1%); HR 1.5 (0.4-6.3) |
| | | | | Week 9 |
| | | | | • Tunneled catheters: 3/178 (1.7%) |
| | | | | • PICCs: 2/183 (1.1%) |
| | | | | Week 9 |
| | | | | • Tunneled catheters: 1/151 (0.7%); HR: 2.0 (0.2-17.7) |
| | | | | • PICCs: 0/125 (0) |
| | | | | Topic-specific outcomes: |
| | | | | Catheter dwell time median, (IQR) |
| | | | | • Tunneled catheters: 24.5 d (14-45) |
| | | | | • PICCs: 11 d (7-18) |
| | | | | • p < 0.001 |
| | | | | Adverse events: NR |

Page **23** of **137**

Author: Shalabi⁵ Year: 2015 Study Design: Retrospective matched cohort study Risk of Bias: Low

Number of patients: N=540

PICC only: N = 180

UVC only: n=180 UVC + PICC: n=180

Setting: tertiary level NICU

Location: Canada

Dates: January 1, 2010 – December 31, 2013

Inclusion Criteria:

- Preterm infants born at less than 30 weeks' gestational age
- Admitted to CNN NICUs within study period
- Received either a UVC or PICC on the first day after birth (day 1) as their venous access
- MATCHING
- Because a small number of infants were expected in the PICC group, eligible infants were first for that group.
- Once the infants in the PICC group were identified, the UVC and UVC + PICC groups were formed by randomly selecting infants from the pool of eligible infants by matching 1:1 for gestational age in weeks, gender, and birth weight 6 100 g.

Exclusion Criteria:

Infants who had a major congenital anomaly

Study Groups: UVC only (n=180)

 Infants who received a UVC on day 1 and did not receive any other central venous access

PICC only (n=180)

 Infants who received a PICC on day 1 and never received a UVC

UVC + PICC (n=180)

 Infants who received a UVC on day 1 that remained in place for a minimum of 4 days followed by placement of a PICC.

Device/agent: Catheter type

Standard preventive measures:

- Patients with multiple episodes of infections were counted once.
- A patient was identified as having a second episode of infection only after 7 days of treatment with the appropriate antibiotic for the previous episode

Outcome Definitions:

CABSI: presence of bacteria or fungus in 1 or more blood cultures obtained from a symptomatic infant after 2 days of placement of a central catheter or within a 48-hour period after catheter removal.

- Did not mandate the need for 2 blood cultures or a blood culture to be drawn from the catheter for diagnosis of CABSI.
- Did not include cultures from the catheter tip in the definition of CABSI
- A patient who had a UVC removed and a PICC inserted on the same day and then developed an infection within 2 days was counted as CABSI associated with UVC and not PICC.

Incidence was calculated per 1000 catheter days and as raw incidence

Rate of any LOS: presence of bacteria or fungus in 1 or more blood cultures from a symptomatic infant

Adverse events: NR

Sampling /Testing strategy: Blood cultures

Other notes:

 Clinical practice of removing UVCs by 5 to 7 days after birth, whereas PICCs are removed mostly when not needed or when complications occur

Primary Outcomes:

CABSI Rate: CABSI / 1000 catheter days

- UVC: 7.8
- PICC: 9.3UVC + PICC: 8.2
- PICC vs UVC: P = 0.60
 - Adj Incident Rate: 1.18 (0.59-2.34)
 - p = 0.64
- PICC vs UVC + PICC: p = 0.55
 - Adj Incident Rate: 1.33 (0.83-2.15)
 - p = 0.23
- UVC vs UVC + PICC: p = 0.89
 - Adj Incident Rate: 1.13 (0.59-2.16)
 - p = 0.71

CABSI Incidence, n (%)

- UVC: 12/180 (7%)
- PICC: 28/180 (15%)
- UVC + PICC: 37/180 (21%)
- PICC vs UVC: P < 0.01
- PICC vs UVC + PICC: p = 0.22
- UVC vs UVC + PICC: p < 0.01

LOS (Late Onset Sepsis)

Rate: / 1000 catheter days

- UVC: 13.7
- PICC: 13.3
- UVC + PICC: 9.3
- PICC vs UVC: P = 0.89
 - Adj Incident Rate: 1.06 (0.64-1.75)
 - p = 0.82
- PICC vs UVC + PICC: p = 0.05
 - Adj Incident Rate: 1.73 (1.15-2.60)
 - p < 0.01
- UVC vs UVC + PICC: p = 0.12
 - Adj Incident Rate: 1.63 (0.97-2.76)
 - p = 0.06

Incidence, n (%)

- UVC: 21/180 (12%)
- PICC: 40/180 (22%)
- UVC + PICC: 42/180 (23%)
- PICC vs UVC: P < 0.01
- PICC vs UVC + PICC: p = 0.80

| Study Information | Population and Setting | Intervention/ Study Groups | Definitions | Results |
|----------------------------|------------------------------|---------------------------------|---|--|
| • | Infants who were moribund | | | • UVC vs UVC + PICC: p < 0.01 |
| | on admission | | | · |
| | Had early onset sepsis | | | Topic-specific outcomes: |
| | Did not receive a central | | | Catheter days |
| | catheter on day 1 | | | • UVC: 1532 days |
| | , | | | • PICC: 3012 days |
| | | | | • UVC + PICC: 4515 days |
| | | | | • p = NA |
| | | | | |
| | | | | Duration of UVC, median (IQR), d |
| | | | | • UVC: 8 (6-10) |
| | | | | PICC: NA |
| | | | | • UVC + PICC: 7 (5-9) |
| | | | | • PICC vs UVC: p = NA |
| | | | | PICC vs UVC + PICC: p = NA |
| | | | | • UVC vs UVC + PICC: p < 0.01 |
| | | | | Duration of PICC, median (IQR), d |
| | | | | • UVC: NA |
| | | | | • PICC: 13 (9-19) |
| | | | | • UVC + PICC: 13 (8-22) |
| | | | | • PICC vs UVC: p = NA |
| | | | | • PICC vs UVC + PICC: p = 0.49 |
| | | | | • UVC vs UVC + PICC: p = NA |
| | | | | |
| | | | | Adverse events: NR |
| Author: Arnts ³ | Number of patients: | Study Groups: | Outcome Definitions: | Primary Outcomes: |
| | N = 232 | UVCs: n=140 UVCs | CLABSI: CDC definition: patients < 1 year | CLABSI: |
| Year: 2014 | Number of lines: | UVCs are typically inserted in | old, laboratory-confirmed bloodstream | Total rate = 20.5 per 1000 CVC days |
| | N= 203 CVCs | the umbilical vein in the first | infection with UVC or PICC in place for a | Total incidence = 13/203 (16.3%) |
| Study Design: | | 2 days postpartum. | minimum of 2 days or in place on the | Incidence: |
| Retrospective | Setting: | | day of event or the day before 4 | • UVC: 21/140 (15%) |
| observational study | Level III NICU | Insertion technique: | | • PICC: 12/63 (19%) |
| D' (D' | | Inserted under aseptic | Laboratory-confirmed BSI: | • p = NR |
| Risk of Bias: | Location: NR | conditions by trained | Criterion 1- one or more positive | |
| Low | . | neonatologists, nurse | blood cultures with the exception of | CDC CLABSI—Laboratory-confirmed BSI |
| | Dates: 16-month period 2005- | practitioners, and | skin micro-organisms, not related to | (Criteria 1 and 2) |
| | 2006 | resident physicians, all of | another source | Total rate = 8 per 1000 CVC days |
| | 1 | whom follow a | Criterion 2- Clinical signs of sepsis | Total incidence = 20/203 (9.8%) |
| | Inclusion Criteria: | standardized protocol | (especially for patients < 1 year old) | Incidence |
| | Gestational age between 24 | outlining the insertion | and two or more positive blood | • UVC: 6/140 (4.3%) |
| | and 42 weeks | practices. | cultures drawn on separate | • PICC: 7/63 (11.1%) |
| | | | occasions with the same micro- | • p = NR |

| Study Information | Population and Setting | Intervention/ Study Groups | Definitions | Results |
|-------------------|---|--|--|---|
| Study Information | Population and Setting CVC (UVC or PICC) inserted in ward Exclusion Criteria: Catheter removed within 24 hours after insertion. CVC inserted in another center. Underwent extracorporeal membrane oxygenation (ECMO) treatment UE | Catheter is fixed with a suture through the umbilical jelly. A second fixation of the catheter with plaster on the abdominal wall using a neo-bridge construction is generally performed for additional safety PICCs: n=63 PICCs inserted via the Seldinger technique. PICCs are inserted by trained neonatologists under maximum aseptic conditions in the NICU. After insertion, the catheter is covered at the insertion site by a sterile transparent film dressing. Device/agent: Catheter site and catheter type Standard preventive measures: The insertion site (not the skin) was disinfected with a 0.5% chlorhexidine/alcohol 70% solution twice daily to conform with hospital policy. The catheter insertion site was examined by trained NICU nurses every 2 hours for signs of inflammation or leakage as a standard of care. The entire drip system for all CVCs was replaced every 96 hours by NICU | organism (including skin microorganisms) and no other infection source Criterion satisfied within a timeframe that did not exceed a gap of 1 day Clinical sepsis: Criterion 3- clinical signs of sepsis (criterion 2) but no or one positive blood culture (only skin microorganisms), with no infection source other than a CVC (in-situ or removed in 24 hours) and a medical reason to initiate sepsis treatment Adverse events: Obstruction: difficulty or inability to flush the catheter or inability to administer fluid in 3 seconds Dislocation: NR Leakage: NR Extravasation/perforation: NR Sampling /Testing strategy: After CVC removal, a tip culture was not routinely performed, except when the CVC was removed due to clinical signs of sepsis. A tip culture was followed by a blood culture when possible. Other notes: NA | Clinical sepsis (Criterion 3): Total rate = 12.4 per 1000 CVC days Total incidence = 20/203 (9.8%) Incidence • UVC: 15/140 (10.7%) • PICC: 5/63 (7.9%) • p = NR Topic-specific outcomes: CVC indwelling time (days): • UVC: 6.9±2.7 • PICC: 10.2±5.2 • p < 0.001 Adverse events Obstruction: • Total rate = 3.1 per 1000 CVC days • Total incidence: 5/203 (2.5%) • UVC: 0/140 (0%) • PICC: 5/63 (7.9%) • p = NR Dislocation: • Total rate = 2.5 per 1000 CVC days • Total incidence: 4/203 (2.0%) • UVC: 4/140 (2.9%) • PICC: 0/63 (0%) • p = NR Leakage: • Total rate = 2.5 per 1000 CVC days • Total incidence: 4/203 (2.0%) • UVC: 4/140 (2.9%) • PICC: 1/63 (1.6%) • p = NR Extravasation/perforation: • Total rate = 1.2 per 1000 CVC days • Total incidence: 2/203 (1.0%) • UVC: 0/140 (0%) |
| 1 | | | | |

| Study Information | Population and Setting | Intervention/ Study Groups | Definitions | Results |
|--|---|---|---|---|
| Author: de Brito ¹⁰ | Population: N= 318 patients | nurses as a standard of care. • All CVCs used were single-lumen CVCs. Study Groups: | Outcome Definitions: | • p = NR Primary Outcomes: |
| Year: 2010 Study Design: Prospective cohort study | N=v461 CVCs Setting: 1 NICU, University Hospital Location: Brazil | UVC: n=33 PICC: n=20 Phlebotomy: n=24 Intracath: n=7 Device/agent: Catheter type | Laboratory-confirmed BSI: isolation of recognized pathogens from blood culture that were not related to infection at another site, with > 38°C fever and with clinical signs of sepsis including apnea, temperature instability, lethargy, feeding | CVC-associated BSI rate/ 1000 catheter days • UVC: 1.7 • PICC: 6.0 • Phlebotomy: 3.5 • Intracath: 1.9 • PICC vs. other catheters: Higher proportion observed in PICC: p<0.01 |
| Risk of Bias: High | Inclusion Criteria: Neonates with at least one CVC placed for >24h, followed up via NHSN. Exclusion Criteria: NR | Standard preventive measures: Catheters removed when no longer required for patient care, when the patient experienced an adverse event, or when catheter exchange was necessary. Catheters removed under aseptic conditions. | intolerance, worsening respiratory distress or hemodynamic instability. Catheter tip colonization: absence of infection signs at the catheter insertion site and microorganism's growth≥103 CFU/mL of the catheter's tips (by quantitative culture). CVC-related BSI: presence of clinical signs for sepsis and positive hemoculture with the same microorganism present on the catheter tip (by quantitative culture) and clinical and microbiological absence of any other source of infection. CVC-associated BSI: bacteremia (isolation of the same organism with identical antibiograms from the blood drawn from peripheral veins and CVC), clinical manifestations sepsis, defervescence after removal of implicated catheter, but without laboratory confirmation of CVC colonization. Incidence density: number of infectious episodes starting during exposure to a specific type of catheter/ number of days of a specific CVC presence times 1000. Sampling /Testing strategy: Blood cultures | CVC-related BSI rate/ 1000 catheter days • UVC: 1.0 • PICC: 0.6 • Phlebotomy: 0.4 • Intracath: 0 Topic-specific outcomes: Dwell time, median, days • UVC: 5.3 • PICC: 13.6 • Phlebotomy: 15.2 • Intracath: 14.8 • UVC vs. other catheters: p = 0.02 Adverse events: NR |
| | | | Other notes: None | |

| Study Information | Population and Setting | Intervention/ Study Groups | Definitions | Results |
|--------------------------------|--|------------------------------------|---|--|
| Author: Chien ¹¹ | Number of patients: | Study Groups: | Outcome Definitions: | Primary Outcomes: |
| | N= 19, 507 | Umbilical venous catheter: n | Nosocomial blood stream infection: one | There was significant variation between |
| Year: 2002 | | = 126 patients | or more positive single organism blood | hospitals in CVC-related infections even after |
| | Number of lines: | Percutaneous catheter: | cultures obtained after 48 h of life in an | adjusting for significant patient characteristics. |
| Study Design: | N = 19,507 | n = 322 patients | infant with clinical suspicion of infection. | a system green signment particular and a second |
| Prospective cohort | , , , , | Tunneled catheter: | To differentiate between nosocomial | Nosocomial BSI: |
| study | Setting: 17 NICUs – Level III | n = 115 patients | and primary (maternal origin) | Incidence: 6.1%; |
| | NICU | | infections, the infant blood culture | Rate: (Incidence/ 1000 Patient Days) |
| Risk of Bias: | | Device/agent: Catheter type | isolates were required to be different | No CVC: 2.9/ 1000 patient days |
| Low | Location: Canada | | from maternal isolates or to occur at | • Crude RR: 1 |
| | | Standard preventive | least 7 days after a treated positive | UVC: 7.2 / 1000 Patient Days |
| | Dates: January 1996 – October | measures: | blood culture obtained during the | • Percutaneous catheter: 13.1 / 1000 Patient |
| | 1997 | NR | first 48 hours of life | <u>,</u> |
| | 1337 | TVIX. | mist 46 hours of me | Days |
| | Inclusion Criteria: | | Infection episode: a positive culture | • Tunneled catheter: 12.1 / 1000 Patient Days |
| | CVC use: umbilical venous | | occurring at least 7 days after a previous | Crude RR |
| | catheter; percutaneously | | treated positive culture or if the culture | • UVC: 2.5 (2.1-3.1) |
| | inserted long catheter or | | isolates were different from the previous | • Percutaneous catheter: 4.6 (4.1-5.3) |
| | , and the second | | culture. | • Tunneled catheter: 4.3 (3.6-5.2) |
| | spaghetti catheter; surgically | | culture. | • p < 0.05 |
| | placed Tunneled catheter. | | At wink any indicate CVC valetad and an annual | |
| | Exclusion Criteria: Viral | | At risk period for CVC-related nosocomial BSI: the period from insertion of a CVC | aRR for BSI: |
| | | | | • UVC: 2.0 (1.7–2.5) |
| | infection | | until removal of CVC or patient | Percutaneous catheter: 3.5 (3.0–4.0) |
| | | | discharge, whichever was shorter. | • Tunneled catheter: 3.1 (2.5–3.8) |
| | | | | |
| | | | Not at-risk period: the length of NICU | Topic-specific outcomes: |
| | | | stay minus the at-risk period. | Median duration of CVC Use (days) |
| | | | | • UVC: 4 ± 8.9 |
| | | | CVC-related nosocomial BSI: All positive | Percutaneous catheter: 10 ± 10.9 |
| | | | blood cultures occurring during the at- | Tunneled catheter: 16 ± 19.1 |
| | | | risk periods | Tumeled catheter. 10 ± 13.1 |
| | | | | Interhospital variation (range) |
| | | | Not CVC-related nosocomial BSI: Positive | • UVC: 1.9% - 60.3% |
| | | | blood cultures occurring during the not | • Percutaneous catheter: 0.2% - 48.1% |
| | | | at-risk periods | |
| | | | Adverse Events | • Tunneled catheter: 0% - 20.5% |
| | | | NR | A diverse accepte |
| | | | | Adverse events |
| | | | Sampling /Testing strategy: Blood | NR |
| | | | cultures | |
| | | | | |
| | | | Other notes: None | |
| Author: Bhandari ¹² | Number of patients: | Study Groups: | Outcome Definitions: | Primary Outcomes: |
| | N=2091 | • UA: n = 1699 | | Nosocomial sepsis: |

| Study Information | Population and Setting | Intervention/ Study Groups | Definitions | Results |
|--------------------|--------------------------------|---|--|---|
| Year: 1997 | Number of lines: | • UV: n = 617 | Nosocomial sepsis: Presence of clinical | Incidence, n (%) |
| | N=2091 CVCs | • CV: n = 294 | signs of infection, initiation of anti- | • UA: 179/1699 (10.5%) |
| Study Design: | | • C: n = 308 | microbial therapy and positive blood | • UV: 81/617 (13.1%) |
| Prospective cohort | Setting: 2 NICUs, 1 University | • PA: n = 189 | cultures obtained from a peripheral | • Tunneled: 99/294 (33.8%) |
| study | Hospital & 1 Regional Hospital | | site or via the catheter after the third | • PC: 96/308 (31.2%) |
| | | Device/agent: Catheter type | postnatal day. | • PAC: 35/189 (18.5%) |
| Risk of Bias: | Location: USA | | _ | • p < 0.0001 |
| Moderate | | Standard preventive | Sampling /Testing strategy: | Incidence by NICU (%) |
| | Dates: | measures: | Blood/catheter tip culture. | • NICU 1: 9.9% |
| | NICU 1: November 11, 1987 - | UA and UV were placed | | • NICU 2: 10.7% |
| | December 31, 1993 | either by the physicians or | Adverse Events: | |
| | | the neonatal nurse | NA | CVC-associated infection incidence, n (%) |
| | NICU 2: January 1, 1989 - | practitioners (NNP) at | | • CV: 17/112 (15.2%) |
| | December 31, 1993 | both NICUs | Other notes: | • PC: 4/79 (5.1%) |
| | | Central venous tunneled | Incidence of infection by comparing | • p < 0.05 |
| | Inclusion Criteria: | catheters (CV) were | different catheter types. | |
| | All neonates admitted to | placed by the same group | To define an association between the | Topic-specific outcomes: (refer to Table 4 for |
| | NICUs during respective | of pediatric surgeons | duration of catheter use, type, and | duration of use by 1-3 days, 4-7 days, 8-14 |
| | study periods | Peripheral arterial | nosocomial sepsis, the incidence of | days, and ≥15 days) |
| | One or more vascular | catheters were placed by | positive blood cultures from time of | Less duration of use highest for UVC |
| | catheters simultaneously | physicians/ NNPs. | insertion of catheter until 3 days after | Greater duration of use highest for UVC and |
| | or sequentially placed | Percutaneous central | removal was analyzed for a | CVC |
| | umbilical artery (UA), | venous placements were | consecutive population subset over | Adverse events: NA |
| | Umbilical venous (UV), | done exclusively by the | 2.5 years at NICU 2 (Jan 7, 91- Dec 31, | |
| | central venous Tunneled | NNPs using a standard | 1993. | |
| | (CV), percutaneously | protocol: sterile technique | | |
| | placed central venous (PC), | and site prep with | | |
| | or peripheral artery (PA). | povidone iodine at both | | |
| | Exclusion Criteria: NR | units. | | |
| | Exclusion Criteria: NK | Catheter maintenance | | |
| | | was done per nursing | | |
| | | protocols at both | | |
| | | hospitals: sterile dressing | | |
| | | and IV tubing changes. | | |
| | | All lines had heparin | | |
| | | infusions. | | |

Page **29** of **137**

Table 16 Risk of Bias of Two Group Studies on Catheter Types

| Author Year | All study groups derived from similar source/reference populations | Attrition not significantly different across study groups | Measure of exposure is valid | Measure of outcome is valid | Investigator blinded or were outcomes well-defined and objective to endpoint assessment | Potential confounders identified | Statistical adjustment for potential confounders done | Funding source(s) disclosed and no obvious conflict of interest | Overall Risk of Bias |
|------------------------------------|--|---|------------------------------|-----------------------------|---|--|--|--|----------------------------|
| Arnts 2014 ³ | ✓ | | ✓ | ✓ | ✓ | ✓ | ✓ | ✓ | Low |
| De Brito 2010 ¹⁰ | ✓ | | ✓ | ✓ | ✓ | ✓ | | | Moderate |
| Bhandari 1997 ¹² | ✓ | | ✓ | ✓ | ✓ | ✓ | | | Moderate |
| Chenoweth 2018 ⁸ | ✓ | ✓ | ✓ | ✓ | | | | ✓ | Moderate |
| Chien 2002 ¹¹ | ✓ | | ✓ | ✓ | √ | ✓ | ✓ | ✓ | Low |
| Geldenhuys 2017 ⁶ | ✓ | | ✓ | ✓ | ✓ | ✓ | ✓ | ✓ | Low |
| Greenburg 2015 ⁷ | ✓ | | ✓ | ✓ | ✓ | ✓ | ✓ | ✓ | Low |
| Konstantinidi 2019 ⁴ | ✓ | ✓ | ✓ | ✓ | | | | ✓ | Moderate |
| Sanderson 2017 ² | ✓ | | ✓ | ✓ | ✓ | ✓ | | ✓ | Low |
| Shalabi 2015 ⁵ | ✓ | | ✓ | √ | ✓ | ✓ | | ✓ | Low |
| Soares 2017 ⁹ | ✓ | NO | ✓ | ✓ | ✓ | ✓ | | ✓ | Low |

C.3. Central Line Insertion Site

Key Question 3: In NICU patients requiring central venous catheters, does the use of one central line catheter insertion site, compared with another, prevent CLABSI?

Table 17 Summary of Findings on Central Line Sites to Prevent CLABSI: PICC Placement in Femoral vs. Non-Femoral Sites

| | | Quantity and Type of Evidence | GRADE of Evidence for Outcome |
|--------------------------|---|---|---|
| Outcome | Findings | and Sample Size | and Limitations of the Evidence |
| Catheter-related sepsis* | • Two observational studies ^{13, 14} conducted in the same NICU population over a slightly different time period found that use of a PICC at a femoral sites was associated with a higher incidence of CRS than at non-femoral sites (N= 518 PICCs) ¹³ (54/240 (22.5%) vs: 34/278 (12.2%); P = 0.002) ¹³ or was a significant risk factor for CRS (10400). ¹⁴ | 2 OBS N= 518 lines ¹³ N= 808 lines ¹⁴ | Very Low ● Imprecision: only one study |

| | | Quantity and Type of | |
|----------------|--|--|---------------------------------|
| | | Evidence | GRADE of Evidence for Outcome |
| Outcome | Findings | and Sample Size | and Limitations of the Evidence |
| | • One observational study ¹⁴ found no difference between groups. | 2 OBS ^{13, 1413, 1413, 1413, 1413,} | Very Low |
| Adverse events | • One observational study ¹³ found that patients with non-femoral central lines were more | 1413, 1413, 1413, 14 | Inconsistency: inconsistent |
| Adverse events | likely to experience phlebitis, catheter site inflammation, or early removal of the central | N= 518 lines ¹³ | results across studies |
| | line. | N= 808 lines ¹⁴ | |

Table 18 Summary of Findings on Central Line Sites to Prevent CLABSI: CVC Placement in Jugular vs. Subclavian vs. Femoral Sites

| Outcome | Findings | Quantity and Type of Evidence and Sample Size | GRADE of Evidence for Outcome and Limitations of the Evidence |
|--------------------------------|--|---|--|
| CLABSI* | One case control study¹⁵ reported a significant increase in the odds of internal jugular placement among NICU patients with CLABSI with internal jugular placements [OR: 2.7 (95% CI: 1.5 – 5.1); p = 0.001], and no difference in the proportion of subclavian, saphenous, external jugular, or brachial placement among NICU patients with CLABSI. One cohort study¹⁶ examining tunneled CVCs reported no difference in the incidence of CLABSI when comparing lines placed in the femoral sites and those placed in the subclavian sites [p = 1.0) | 2 OBS n = 179 lines ¹⁵ n = 601 lines ¹⁶ | Low |
| Catheter-associated Infection* | One observational study¹⁷ found that the use of subclavian sites was associated with a lower rate of catheter-associated infections compared with the jugular vein for implanted catheters in NICU patients with surgically-implanted CVCs. (p<0.01). | 1 OBS n = 236 lines ¹⁷ | Very Low ■ Imprecision: only one study |
| Catheter-related sepsis* | • One observational study ¹⁸ found that the use of femoral sites was associated with a lower rate of catheter-related sepsis when compared with sites in the neck including jugular and subclavian sites for long-term, tunneled catheters in NICU patients. (p = 0.032). | 1 OBS n = 137 lines ¹⁸ | Very LowImprecision: only one studyStudy Quality: study at high risk of bias |

Table 19 Summary of Findings on the Efficacy of Central Line Site to Prevent CLABSI: CVC Placement in Upper vs. Lower Extremities

| Outcome | Findings | Quantity and Type of Evidence and Sample Size | GRADE of Evidence for Outcome and Limitations of the Evidence |
|-----------------------|--|--|---|
| CLABSI* | • Two cohort studies ^{19, 20} reported no significant difference in CLABSI incidence or rates between insertion sites (Adjusted OR: 1.23 (95% CI: 0.58-2.60); p = 0.57) ¹⁹ or [p = 0.941]. ²⁰ | 3 OBS n = 1,104 lines ¹⁹ n = 365 lines ²⁰ n = 179 lines ¹⁵ | Low |
| Catheter related-BSI* | • One observational study ²¹ reported no significant difference in CRBSI incidence between insertion sites (UE: 43/370 (11.6%) vs LE: 10/107 (9.3%)). | 1 OBS n = 477 lines ²¹ | Very Low • Imprecision: only one study |
| Sepsis* | One observational study²⁰ reported no difference in the proportion of sepsis for PICCs inserted in upper and lower extremities in NICU patients (p = 0.941) | 1 OBS N= 365 lines ²⁰ | Very Low • Imprecision: only one study |

| | | Quantity and Type of Evidence | GRADE of Evidence for Outcome |
|------------------|---|--|--|
| Outcome | Findings | and Sample Size | and Limitations of the Evidence |
| Presumed Sepsis* | • One observational study ²² reported no significant difference between insertion sites (UE: 31 (8.3) vs LE: 18 (7.1) p = 0.6006). | 1 OBS n = 626 lines ²² | Very Low • Imprecision: only one study |
| Adverse Events | No significant difference was reported between groups for thrombus,²⁰ phlebitis,^{19, 21, 22} | 4 OBS n = 1,104 lines ¹⁹ n = 477 lines ²¹ n = 626 lines ²² N= 365 lines ²⁰ | Low |

Table 20 Extracted Information on Central Line Sites

| Study Information | Population and Setting | Intervention/ Study Groups | Definitions | Results |
|--|--|--|---|--|
| Author: Elmekkawi ²⁰ | Number of patients: | Study Groups: | Outcome Definitions: | Sepsis during the line: |
| | N = 365 | UE PICCS: n=138 | Sepsis during the line: blood culture taken | Incidence, n (%) |
| /ear: 2019 | Number of lines: | Via basilic, cephalic, median | a minimum of 24 hours after catheter | • UE: 18/138 (13.0%) |
| | N=365 PICC lines | cubital, or axillary veins | insertion and a maximum of 48 hours | • LE: 29/227 (12.8%) |
| Study Design: | Setting: NICU at | LE PICCs: n=227 | after catheter removal was positive | • p = 0.941 |
| Retrospective | quaternary children's | Via greater saphenous vein, | | Coagulase-negative staphylococcus incidence, n |
| cohort | hospital | lesser saphenous vein, | Adverse events: | (%) |
| | | dorsal venous arch, or | Mortality: death | • UE: 12/138 (8.7%) |
| Risk of Bias: Low | Location: Canada | popliteal vein | Mechanical: occlusion or leaking | • LE: 17/227 (7.5%) |
| | | | Interstitial: NR | , , , |
| | Dates: January 2005 – | Device/agent: Catheter site | Pleural or pericardial effusion: NR | S. aureus incidence, n (%) |
| | August 2010 | | Phlebitis: NR | • UE: 1/138 (0.7%) |
| | | Standard preventive | Thrombus: NR | • LE: 1/227 (0.4%) |
| | Inclusion Criteria: | measures: | | Group B streptococcus incidence, n (%) |
| | Neonates who had | Majority of PICCs were | Sampling /Testing strategy: Blood | • UE: 0/138 (0%) |
| | PICC lines placed in | inserted by specialized | cultures | • LE: 1/227 (0.4%) |
| | the NICU | PICC nurses | | Enterococcus incidence, n (%) |
| | | Catheter choice and | Other notes: None | • UE: 0/138 (0%) |
| | Exclusion Criteria: | insertion site were | | • LE: 1/227 (0.4%) |
| | Lines inserted by | guided by operator | | Klebsiella incidence, n (%) |
| | interventional | preference and vein | | • UE: 1/138 (0.7%) |
| | radiology | availability | | • LE: 3/227 (1.3%) |
| | Patients that were | Procedure was | | E. coli incidence, n (%) |
| | transferred out of the | performed at the | | , , , |
| | NICU with a PICC in | bedside and ultrasound | | • UE: 2/138 (1.4%) |
| | situ, or died with a | guidance was not used | | • LE: 1/227 (0.4%) |
| | line <i>in situ</i> | Post insertion X-rays | | Enterobacter incidence, n(%) |
| | PICCS that | were taken with the | | • UE: 1/138 (0.7%) |
| | were malpositioned on | shoulder abducted at 30 | | • LE: 2/227 (0.9%) |
| | the insertion X-ray | degrees for UE PICCs and | | S. marcescens incidence, n (%) |

| Study Information | Population and Setting | Intervention/ Study Groups | Definitions | Results |
|------------------------------|--|---|---|--|
| | that could not be used | the hips in 'frog' position | | • UE: 0/138 (0%) |
| | for infusion and | for LE PICCs | | • LE: 2/227 (0.9%) |
| | removed immediately | A repeat X-ray to | | Proteus incidence, n (%) |
| | post X-ray | confirm final tip position | | • UE: 1/138 (0.7%) |
| | PICCs removed within | was done if the catheter | | • LE: 0/227 (0%) |
| | 24 hours of insertion | was pulled by more than | | (|
| | for malposition | 1 cm | | Topic-specific outcomes: |
| | | The routine unit practice | | Duration of catheter median, days (IQR) |
| | | was to remove non- | | • UE: 17 days (8-27) |
| | | central PICCs within 24 | | • LE: 16 days (9-30) |
| | | hours of insertion | | ==: == uays (5 55) |
| | | | | Adverse events |
| | | | | Mortality, n (%) |
| | | | | • UE: 7/138 (5.1%) |
| | | | | • LE: 14/227 (6.2%) |
| | | | | • p = 0.818 |
| | | | | Mechanical (occlusion or leaking), n (%) |
| | | | | • UE: 14/138 (10.1%) |
| | | | | • LE: 28/227 (12.3%) |
| | | | | Interstitial, n (%) |
| | | | | • UE: 3/138 (2.2%) |
| | | | | • LE: 3/227 (1.3%) |
| | | | | Pleural or pericardial effusion, n (%) |
| | | | | • UE: 3/138 (2.2%) |
| | | | | , , , |
| | | | | • LE: 0/227 (0%) |
| | | | | Phlebitis, n (%) |
| | | | | • UE: 1/138 (0.7%) |
| | | | | • LE: 10/227 (4.4%) |
| | | | | Thrombus, n (%) |
| | | | | • UE: 0/138 (0%) |
| | | | | • LE: 1/227 (0.4%) |
| Author: Garcia ¹⁵ | Number of patients: | Case: | Outcome Definitions: | Primary Outcomes: |
| V 2010 | N = 179 patients | CLABSI: n=74 | CLABSI: CDC 2018 definition | Placement site of CVC: |
| Year: 2019 | Number of lines: | Control | Patient ≤1 year of age has at least one | Internal jugular, n/N (%) |
| Study | N=179 lines | Control: Non-CLABSI: n=105 | of the following signs or symptoms: | • OR: 2.7 (95% CI: 1.5-5.1); P = 0.001 |
| Study Design: Nested case- | Setting: | NUII-CLADSI. II=1US | fever (>38.0°C), hypothermia | • Case: 43/74 (58.1%) |
| control | Third-care level NICU | Device/agent: Catheter site; | (<36.0°C), apnea, or bradycardia, and | • Control: 35/105 (33.3%) |
| CONTROL | THIRD COLC ICVELINICO | double lumen catheter | Organism(s) identified in blood is (are) | • p = 0.001 |
| Risk of Bias: Low | Location: Mexico | double fullett catheter | not related to an infection at another | Subclavian (percutaneous insertion), n/N (%) |
| 01 51631 2011 | | | site, and | • Case: 17/74 (23%) |

| | | Intervention/ Study Groups | Definitions | Results |
|----------------------------|---|----------------------------------|--|---|
| | Dates: January 2014 – December 2015 Inclusion Criteria: • Patients with installation of a CVC during their hospital stay at the NICU were included • Patients with first CVC installation and those with CVC duration ≥48 hours • Cases were neonates diagnosed with CLABSI • Controls were those neonates with a CVC during the same period but who did not develop a CLABSI Exclusion Criteria: Patients who had a catheter installed in another hospital | Standard preventive measures: NR | The same common commensal is identified by a culture or non-culture based microbiologic testing method, from two or more blood specimens collected on separate occasions Adverse events: CLABSI-related mortality: a death directly related to the infection which occurred during active infection event and no other underlying cause of fatal outcome was present Sampling /Testing strategy: Two-set of blood cultures were obtained in patients with a suspected infection Disinfection with 2% iodine-povidone were performed One peripheral blood culture was obtained along with a catheter-drawn blood culture Other notes: None | Control: 27/105 (25.7%) p = 0.67 Saphenous, n/N (%) Case: 7/74 (9.5%) Control: 16/105 (15.2%) p = 0.25 External jugular, n/N (%) Case: 4/74 (5.4%) Control: 7/105 (6.7%) p = 0.98 Upper limb, n/N (%) Case: 1/74 (1.3%) Control: 12/105 (11.4%) p = 0.01 Brachial, n/N (%) Case: 1/74 (1.3%) Control: 5/105 (4.8%) p = 0.21 Lower limb, n/N (%) Case: 1/74 (1.3%) Control: 3/105 (2.8%) p = 0.64 Double-lumen catheter: OR: 10.0 (95% CI: 2.3-44.3); P = 0.0001 Case: 72/74 (97.3%) Control: 82/105 (78.1%) Topic-specific outcomes: CVC indwelling total time >21 days, n/N (%): OR: 2.9 (95% CI: 1.5-5.4); P = 0.001 Case: 37/74 (50.0%) Control: 27/105 (25.7%) Adverse events CLABSI-related mortality, n/N (%) Case: 5/74 (6.8%) |
| Author: Litz ¹⁶ | Number of patients: N = 601 | Study Groups: T-CVC: n=134 | Outcome Definitions: CLABSI: CDC 2015 definition | Control: NR Primary Outcomes: CLABSI |

| Study Information | Population and Setting | Intervention/ Study Groups | Definitions | Results |
|-------------------|---------------------------------|--|--|---|
| Year: 2017 | Number of lines: | PICC: n=467 | | Incidence, n/N (%): |
| | N=601 lines | | Line utilization ratio: the number of | • T-CVC: 14/134 (10.2%) |
| Study Design: | | Device/agent: Catheter type | central line days divided by the number of | • PICC: 10/467 (2.1%) |
| Retrospective | Setting: NICU | and site | patient days | • p = NR |
| cohort | | | | Incidence, % |
| | Location: USA | Standard preventive | Adverse events: | T-CVC placed in femoral or saphenous |
| Risk of Bias: Low | | measures: | Line complications: mechanical (broke, | vein: 8.5% |
| | Dates: November | PICC lines are the | infiltrated occluded), local concerns | T-CVC placed in subclavian or jugular vein: |
| | 2008 – October 2015 | preferred modality of | (erythema, swelling, phlebitis), | 10.8% |
| | In about an Orithauta. | vascular access in | malposition/ migration, or other (pleural | • p = 1.0 |
| | Inclusion Criteria: | neonates and T-CVCs are | effusion, arrhythmia, deep venous | Incidence, rate/ 1000 line days |
| | Patients in the NICU | typically placed in long- | thrombosis) | • OR: 0.50 (95% CI: 0.11-2.22); P = 0.55 |
| | who had T-CVCs | term access is needed or | Sampling /Tosting strategy | - |
| | placed between | alternative vascular | Sampling /Testing strategy: | • In use T-CVC: 2.2 |
| | November | access is unable to be | • NR | • Idle T-CVC: 1.1 |
| | 2008 – October 2015 | obtained | Other notes: None | • p = NR |
| | or PICCs placed between July | PICCs are placed and | Other notes: None | Incidence, rate/ 1000 line days |
| | 20014 – October 2015 | removed by a dedicated | | • OR: 0.50 (95% CI: 0.11-2.22); P = 0.55 |
| | 20014 - October 2013 | NICU vascular access | | • In use PICC: 1.3 |
| | Exclusion Criteria: | team comprised of trained nurses, nurse | | • Idle PICC: 0 |
| | Patients who died or were | practitioners, and | | • p = NR |
| | discharged with a central | physicians | | |
| | venous catheter and | T-CVCs are placed by | | Topic-specific outcomes: |
| | those who were not | surgeons and removed | | Line utilization ratio |
| | yet discharged were | by surgical nurse | | • T-CVC: 0.52 |
| | excluded | practitioners, fellows, or | | • PICC: 0.27 |
| | | attendings | | • p < 0.001 |
| | | Daily chlorhexidine | | · |
| | | gluconate treatments | | Adverse events |
| | | for patients >36 weeks | | Line complications, n/N (%) |
| | | and >1000g | | • T-CVC: 9/134 (6%) |
| | | Routine tubing and | | • PICC: 32/467 (6.8%) |
| | | sterile cap changed | | • p = NR |
| | | every 96 hours or 24 | | F |
| | | hours for lines running | | |
| | | lipids, propofol, or blood | | |
| | | products | | |
| | | Heparinized intravenous | | |
| | | fluid at a minimal rate | | |
| | | (1ml/h) to maintain | | |
| | | patency in idle lines | | |
| | | | | |

| Study Information | Population and Setting | Intervention/ Study Groups | Definitions | Results |
|---------------------------------|--|---|---|---|
| | | Daily discussion of the | | |
| | | need for a central line on | | |
| | | rounds | | |
| Author: Bashir | Number of patients: | Study Groups: | Outcome Definitions: | Primary Outcomes: |
| | N = 827 patients | UE PICCs: n=593 | CLABSI: (CDC) | CLABSI: |
| Year: 2016 ¹⁹ | Number of lines: | Via cephalic and basilica | Confirmed primary bloodstream | aOR: 1.23 (95% CI: 0.58-2.60); P = 0.57 |
| | N=1104 PICC lines | veins | infection with one of following clinical | Rate/ 1000-line days |
| Study Design: | | LE PICCs: n=234 | signs of infection (fever, hypothermia, | ● UE: 4.7 |
| Retrospective cohort | Setting: | Via saphenous veins | apnea, or bradycardia) | • LE: 3.3 |
| study | Tertiary NICU | | Presence of central catheter at the | • p = NR |
| | | Device/agent: Catheter site | time of or within 48 hours before the | Incidence, n (%) |
| Risk of Bias: | Location: Canada | | onset of the infection | • UE: 35/593 (5.9%) |
| Low | | Standard preventive | | • LE: 10/234 (4.2%) |
| | Dates: January 1, 2006 – | measures: | Incidence of CLABSI: infection episodes | • p = 0.35 |
| | December 31, 2010 | Data from first time PICC | per 1000 catheter days | F - 2 - 2 |
| | | used if more than one | | Topic-specific outcomes: |
| | Inclusion Criteria: | PICC placed during | Adverse events: | Duration of catheter median, days (IQR) |
| | All preterm infants | hospital stay | Mechanical complications considered | • UE: 10 days (6-15) |
| | (age < 37 complete | PICC lines were placed at | present if there was a line infiltration, | • LE: 10.5 days (5-17) |
| | weeks) | the baby's bedside, | occlusion, phlebitis, and dislodgement, | • p = 0.81 |
| | 1st time PICCs inserted | under sterile conditions, | resulting in removal of PICC | p 0.02 |
| | during study period | by a dedicated team of | Line infiltration: extravasation of fluid | Adverse events |
| | | transport nurses, | into soft tissue around the region of | Infiltration, n (%) |
| | Exclusion Criteria: | neonatal physicians, and | the catheter tip. | • UE: 89/593 (15%) |
| | Infants with | nurse practitioners | Line occlusion: inability to infuse fluid | • LE: 15/234 (6.4%) |
| | incomplete PICC data | Site of insertion was | Phlebitis: presence of a linear red | • p = 0.001 |
| | PICCs inserted from | selected at the | streak developing along the superficial | UE vs LE, n (%) |
| | sites other than upper | discretion of the inserter | veins from the catheter insertion site. | • Right: 56/320 (17.5%) vs 14/152 (9.2%) |
| | or lower extremity | based on the | Dislodgement: NR | • Left: 33/273 (12%) vs 1/82 (1.2%) |
| | Neonates who were | accessibility of veins. | | • p < 0.001 |
| | transferred out to | During the study period, | Sampling /Testing strategy: | Adjusted OR: 2.41 (95% CI: 1.36-4.29); P = 0.003 |
| | other hospitals with an | single lumen catheter | Blood cultures | Occlusion, n (%) |
| | indwelling catheter | 20-30 cm long with an | | • UE: 52/593 (8.7%) |
| | and who did not | introducer cannulae. | Other notes: NA | |
| | return the final data | After the catheter was | | • LE: 31/234 (13.2%) |
| | | inserted, catheter tip | | • p = 0.054 |
| | | position was confirmed | | UE vs LE, n (%) |
| | | by radiograph with the | | • Right: 21/320 (6.5%) vs 23/152 (15.1%) |
| | | limbs in standard resting | | • Left: 31/273 (11.3%) vs 8/82 (9.7%) |
| | | position, and repeat | | • p = 0.02 |
| | | radiographs were taken | | • Adjusted OR: 0.68 (95% CI: 0.41-1.10); P = 0.12 |
| | | if there was a | | Phlebitis, n (%) |
| | | manipulation. | | • UE: 21/593 (3.5%) |
| | | | | • LE: 9/234 (3.8%) |

| Study Information | Population and Setting | Intervention/ Study Groups | Definitions | Results |
|---------------------------------|--------------------------|--|--|--|
| | | Optimal placement for | | • p = 0.83 |
| | | UE: catheter tip lying | | UE vs LE, n (%) |
| | | beyond midclavicular | | • Right: 12/320 (3.7%) vs 6/152 (3.9%) |
| | | area and up to 1 cm at | | • Left: 9/273 (3.3%) vs 3/82 (3.6%) |
| | | the junction of right | | • p = 0.98 |
| | | atrium and superior | | Adjusted OR: 0.88 (95% CI: 0.39-1.98); P = 0.76 |
| | | vena cava | | Dislodgement incidence, n (%) |
| | | Optimal placement for | | • UE: 1/593 (0.1%) |
| | | LE: catheter tip located | | • LE: 0/234 (0%) |
| | | in the inferior vena cava | | • p = 0.63 |
| | | below the diaphragm | | UE vs LE incidence, n (%) |
| | | Heparin was infused in | | • Right: 1/320 (0.31%) vs 0/152 (0%) |
| | | all PICCs as per standard | | • Left: 0/273 (0%) vs 0/82 (0%) |
| | | unit policy. | | • p = 0.66 |
| | | All catheters were | | - p = 0.00 |
| | | removed either after | | |
| | | completion of | | |
| | | intravenous therapy or | | |
| | | prematurely if they | | |
| | | developed | | |
| | | complications. | | |
| Author: | Number of patients: | Study Groups: | Outcome Definitions: | Primary Outcomes: |
| Wrightson | N = 559 | Upper extremities | CLABSI: CDC definition | CLABSI: |
| Year: 2013 ²² | Number of lines: | N=374 PICCs (59.7%) | Presumed sepsis: collective term for PICCs | CLABSI incidence/ PICCs removed for presumed |
| | N= 626 PICCs | For an upper extremity vein, | removed for suspected sepsis or positive | sepsis: 28/50 (56%) |
| Study Design: | | the ideal tip location is in | blood cultures | CLABSI Rate for PICCs removed because of |
| Retrospective cohort | After Exclusion: | the superior vena cava at | | confirmed sepsis: 2.86/ 1000 catheter days |
| | N = 528 patients | T2-T4 resting just above the | Adverse Events: | Presumed sepsis, n (%) |
| Risk of Bias: | N = 655 PICCs | right atrium. (NANN PICC | Nonelective removal: unresolvable PICC | • Incidence: 50/626 (8%) |
| Low | Excluded n=29 | guidelines) | complication leading to removal of the | • UE: 31 (8.3) |
| | | • Axillary 62 (16.6%) | PICC prior to the completion of therapy | • LE: 18 (7.1) |
| | Setting: Level III NICU | Basilic 119 (31.8) | for which the PICC was initially placed | • p = 0.6006 |
| | | • Cephalic 186 (49.7%) | (leaking, clotting, presumed sepsis, | PICCs removed for any complication |
| | Location: USA | Unspecified 7 (1.9%) | positive blood cultures, catheter | Central Tip vs Non-central Tip |
| | | | contamination, thrombosis, edema, | • UE: 73 (72%) vs 29 (28%) |
| | Dates: January 1, 2004 – | Lower extremities | phlebitis, pleural effusion, cardiac | • p = 0.0001 |
| | December 31, 2009 | N=252 PICCs (40.3%) | tamponade, central tip required, broken | • LE: 50 (94%) vs 3(6%) |
| | | For lower extremity veins, | catheter, dislodgement, or malposition.) | • p = 0.7 |
| | Inclusion Criteria: | the tip should be in the | a | · |
| | All PICCs placed in the | inferior vena cava (IVC) at | Clotted: NR | Topic-specific outcomes: |
| | NICU during the | the level of the diaphragm, | | PICC dwell time, range (mean ± SD; median): |
| | timeframe | outside the heart. (NANN | Leaking: NR | • UE: 0-160 days (15 ± 13; 13) |
| | | PICC guidelines) | - 1 | • LE: 0-76 days (16 ± 11.6; 13.5) |
| | | | Edema/infiltrated: NR | Page 37 of 13 |

| Central and non-central veins Exclusion Criteria: Incomplete data | Study Information | Population and Setting | Intervention/ Study Groups | Definitions | Results |
|--|-------------------|-------------------------|--|---|---|
| Exclusion Criteria: Incomplete data Neonate transfer with the PICC indwelling Neonate transfer with the PICCs indwelling Neonate transfer with the PICCs indeed in the clinical nurse speciality trained nurses specially trained nurses specially trained nurses has inserted and maintained PICCs at the study hospital NICU since 1999. On rare occasions, when a PICC team inserter was not available or was unsuccessful at the insertion, PICCs were placed by a physician. Author: Number of patients: Tsai N = 534 Number of lines: N = 680 Revecutaneously inserted CVCs Study Design: Retrospective cohort study Location: Taiwan Non-femoral n = 278 (190 patients) Low Oates: January 2004 — December 2007 D | | Central and non- | | | • p = 0.2038 |
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| • Incomplete data • Neonate transfer with the PICC indwelling • None of the study infants had concurrent PICCs • Under the supervision of the neonatologists and the clinical nurse specialist, a team of specially trained nurses has inserted and maintained PICCs at the study hospital NICU since 1999. On rare occasions, when a PICC team inserter was not available or was unsuccessful at the insertion, PICCs were placed by a physician. Author: Tsai Author: Number of patients: N = 534 Number of lines: N = 808 Percutaneously inserted CVCs Study Design: Study Design: Study Design: Study Design: Costing: Level III NICU Location: Taiwan Dates: January 2004 – December 2007 Location: Taiwan Premature infants with BW ≤ 1500g Dates: January 2004 – December 2007 Location: Taiwan Premature infants with BW ≤ 1500g Premature infants with BW ≤ 1500g Non-femoral n = 278 (190 patients) Location: Taiwan Premature infants with BW ≤ 1500g Premature infants with BW ≤ 1500g Non-femoral n = 278 (190 patients) Location: Taiwan Premature infants with BW ≤ 1500g Premature red sterak developed along the superficial veins from the information and phlebitis categorized as philebitis Premature infants with BW ≤ 1500g Premature red sterak developed along the superficial veins from the catheter-related sepsis (and the clurte negative; patients with both inflammation and phlebitis categorized as philebitis Premature infants with BW ≤ 1500g Premature of the study directly to a neonate's death. 2-2% chlorhexidine gluconate for skin antistepsis was implemented during the study period. Authors do not note when, and note it was only for infants weighing > 1200 g or older than 2 weeks. Authors note "its impact on too the sepsis rates during the study period is unknown." Prica (4.1.1) Prica (1.1) Prica | | | | | Adverse events |
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| the PICC indwelling The PICC indwelling Infants had concurrent PICCs | | Incomplete data | measures: | No PICC complications contributed | • UE: 4 (1.1) |
| PICCS • Under the supervision of the neonatologists and the clinical nurse speciality, a team of specially trained nurses has inserted and maintained PICCS at the study hospital NICU since 1999. On rare occasions, when a PICC team inserter was not available or was unsuccessful at the insertion, PICCS were placed by a physician. Author: Number of patients: N = 534 Number of lines: Percutaneously inserted CVCS Study Design: Study Design: Study Design: Risk of Blas: Location: Taiwan Dates: January 2004 – December 2007 Inclusion Criteria: • Premature infants with BW × 1500g PICS • Under the supervision of the supervision of the neonatologists and the clinical nurse specially trained nurse specially, ta team of specially trained nurses has inserted and the clinical nurse specially trained nurses when, and note it was only for infants weighing >1200 g or older than 2 weeks. Authors note it was only for infants weighing >1200 g or older than 2 weeks. Authors note it was only for infants weighing >1200 g or older than 2 weeks. Authors note it with sepsis rates during the study period. Authors do not note when, and note it was only for infants weighing >1200 g or older than 2 weeks. Authors note it with sepsis rates during the study period. Authors do not note when, and note it was only for infants weighing >1200 g or older than 2 weeks. Authors note it wis only for infants weighing >1200 g or older than 2 weeks. Authors note it wis only for infants weighing >1200 g or older than 2 weeks. Authors note it wis only for infants weighing >1200 g or older than 2 weeks. Authors note it wis only for infants weighing >1200 g or older than 2 weeks. Authors note it wis only for infants weighing >1200 g or older than 2 weighing >1200 g or older than 2 weeks. Authors note it wis only for infants weighing >1200 g or older than 2 weeks. Authors note it wis only for infants weighing >1200 g or older than 2 weeks. Authors note it wis only for infants weighing >1200 g or older than 2 weeks. Authors note. It is fe. 4. • Leak (1 | | Neonate transfer with | None of the study | directly to a neonate's death. | • LE: 5 (2) |
| Inclusion Criteria: Punder the supervision of the neonatologists and the clinical nurse specialist, a team of specially trained nurses has inserted and maintained PICCs at the study heriod is unknown." Author: Number of patients: Tsai N = 534 Number of lines: Percutaneously inserted CVCs 2011 ²⁴ | | the PICC indwelling | infants had concurrent | • 2% chlorhexidine gluconate for skin | • p = 0.4958 |
| the neonatologists and the clinical nurse specialist, a team of specially trained nurses has inserted and maintained PICCs at the study hospital NICU since 1990, no rate occasions, when a PICC team insertier was not available or was unsuccessful at the insertion, PICCs were placed by a physician. Author: Tsai Author: Tsai N = 534 Number of lines: Year: Year: Year: Year: Year: Study Bosign: Study Design: Study Design: Retrospective cohort study Dates: January 2004 – December 2007 Risk of Bias: Low Dates: January 2004 – December 2007 Bigs of Bias: Low Dates: January 2004 – December 2007 Dates: January 2004 – December 2007 Inclusion Criteria: Premature infants with BW s 1500g When, and note it was only for infants weighing 1200 g or older than 2 weeks. Authors note "its impact on the sepsis rates during the study period is unknown." When, and note it was only for infants weighing 1200 g or older than 2 weeks. Authors note "its impact on the sepsis rates during the study period is unknown." Weeks, Authors note "its impact on the sepsis rates during the study period is unknown." Weeks, Authors note "its impact on the sepsis rates during the study period is unknown." **UE: 16 (6.4) **p = 0.05976 Let: 4 (1.6) **p = 0.0605 Edema/, Infiltrated, n (%) **UE: 16 (4.3) **LE: 4 (1.6) **p = 0.01574 **Outcomes: Catheter-related sepsis (CRS): culture confirmed; at least 1 positive culture of blood obtained from a peripheral vein, clinical features consistent with bloodstream infection, no other site of infection, and a PICC in place for at least 3 days. Adverse events: Phlebitis: a linear red streak developed along the superficial veines from the insertion site; can be culture negative; patients with both inflammation and phlebitis at elements with both inflammation and phlebitis at elements with both inflammation and phlebitis at elements and phlebitis at elements and phlebitis. **Premature infants with both inflammation and phlebitis at elements and phlebitis at elements and phlebitis at elements and | | | PICCs | antisepsis was implemented during | Clotted, n (%) |
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| the clinical nurse specialist, a team of specially trained nurses has inserted and maintained PICCs at the study hospital NICU since 1999. On rare occasions, when a PICC team inserter was not available or was unsuccessful at the insertion, PICCs were placed by a physician. Author: Tsai Author: Number of patients: N = 534 Number of lines: Percutaneously inserted CVCs 1011 ¹⁴ Netrospective cohort study Study Design: Retrospective cohort study Study Retrospective cohort study Study Permature infants with BW ≤ 1500g The clinical nurse specialist, a team of specialist, a team of specialist, a team of specialist, a team of the sepsis rates during the study period is unknown." Wegks. Authors note "its impact on the sepsis rates during the study period is unknown." Wegks. Authors in the study period is unknown." **Durble of patients** Number of patients: N = 534 Number of lines: Percutaneously inserted CVCs (334 patients) Non-femoral n = 278 (190 patients) Non-femoral n = 278 (190 patients) Setting: Level III NICU Since 1999. On rare occasions, when a PICC team inserter was not available or was unsuccessful at the insertion special period is unknown." **Durble of lines: Percutaneously inserted complications: Catheter-related sepsis (CRS): culture confirmed; at least 1 positive culture of blood obtained from a peripheral vein, clinical features consistent with bloodstream infection, no other site of infection, and a PICC in place for at least 3 days. Adverse events: Phebitis: a linear red streak developed along the superficial veins from the insertion site; can be culture negative; patients with both inflammation and phlebitis categorized as phlebitis • p = 0.5976 Leaking, 1 (%) • UE: 15 (4) • UE: 15 (4) • UE: 15 (4) • UE: 5 (2) • p = 0.1574 **Catheter-related complications: 271/534 patients experienced 368 total catheter-related sepsis (Incidence: 134/368 (36.4%) • New Percutaneous CVC: 8.8 cases per catheter-days • New Percutaneous CVC: 8.8 cases per catheter-days • New Percutaneous CVC: 9.9 cases per cat | | | the neonatologists and | when, and note it was only for infants | |
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| specially trained nurses has inserted and maintained PICCs at the study hospital NICU since 1999. On rare occasions, when a PICC team inserter was not available or was unsuccessful at the insertion, PICCs were placed by a physician. Author: Tsai N=534 Number of lines: N=534 Number of lines: N=608 Percutaneously inserted CVCs (234 patients) Non-femoral n= 278 (190 patients) Study Design: Setting: Level III NICU Study Group: CVCs (334 patients) Setting: Level III NICU Setting: Level III NICU Location: Taiwan Risk of Bias: Low Dates: January 2004 – December 2007 December 2007 Low Defaulte inserted CVCs used before June 2006— single lumen silicone catheter with an introduction cannula New settions is catheter with an introduction cannula New settion site; can be culture negative; patients with both inflammation and phlebits categorized as phlebits New Percutaneous VCC: 9.9 cases per catheter-days New Percutaneous VCC: 9.9 cases per catheter-days New Percuta | | | specialist, a team of | weeks. Authors note "its impact on | · · |
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| since 1999, On rare occasions, when a PICC team inserter was not available or was unsuccessful at the insertion, PICCs were placed by a physician. Author: Tsai N=534 Number of lines: N= 808 Percutaneously inserted CVCs Study Design: Setting: Level III NICU Study Design: Study Design: Study Location: Taiwan Sinfants) Dates: January 2004 − Low December 2007 Inclusion Criteria: Premature infants with BW ≤ 1500g Since 1999, On rare occasions, when a PICC team inserter vas not available or was unsuccessful at the insertion piCCs were placed by a physician. Outcome Definitions: Catheter-related sepsis (CRS): culture confirmed; at least 1 positive culture of blood obtained from a peripheral vein, clinical features consistent with bloodstream infection, no other site of infection, and a PICC in place for at least 3 days. Adverse events: Phemature infants with BW ≤ 1500g Inclusion Criteria: Premature infants with BW ≤ 1500g Author: Outcome Definitions: Catheter-related sepsis (CRS): culture confirmed; at least 1 positive culture of blood obtained from a peripheral vein, clinical features consistent with bloodstream infection, no other site of infection, and a PICC in place for at least 3 days. Adverse events: Phemature infants with BW ≤ 1500g Adverse events: Primary Outcomes: Catheter-related sepsis (CRS): culture confirmed; at least 1 positive culture of blood obtained from a peripheral vein, clinical features consistent with bloodstream infection, no other site of infection, and a PICC in place for at least 3 days. Adverse events: Phemature infants with blood brained from a peripheral vein, clinical features consistent with bloodstream infection, no other site of infection, and a PICC in place for at least 3 days. Catheter-related sepsis Incidence: 134/368 (36.4%) Po 0.680 Risk of Bias: Date: 15 (4) Location: Taiwan Outcome Definitions: Catheter-related sepsis (CRS): culture catheter-related sepsis (Incidence: 134/368 (36.4%) Po 0.680 Rist of Bias: Date: 15 (4) Outcome Definitions: Catheter-related se | | | ' ' | | • |
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| Number of lines: N=808 Percutaneously inserted CVCs (334 patients) Study Design: Retrospective cohort study Coation: Taiwan Dates: January 2004 − December 2007 Inclusion Criteria: Percutaneously inserted CVCs (334 patients) Percutaneously inserted CVCs (334 patients) CVCs (334 patients) Non-femoral n= 278 (190 patients) Popatients Percutaneously inserted CVCs (334 patients) Non-femoral n= 278 (190 patients) Popatients Percutaneously inserted CVCs (334 patients) Non-femoral n= 278 (190 patients) Permoral n= 278 (190 patients) Permoral n= 278 (190 patients) Permoral n= 240 (183 infants) Adverse events: Phebitis: a linear red streak developed along the superficial veins from the insertion site; can be culture negative; patients with both inflammation and phlebitis categorized as phlebitis Percutaneously inserted CVCs (334 patients) Catheter-related sepsis Incidence: 134/368 (36.4%) Old Peripheral CVC: 46/290 (15.9%) Per 0.680 Rate Old Percutaneous CVC: 8.8 cases per 10 catheter-days New Percutaneous CVC: 9.9 cases per 10 catheter-days | | • | ' ' | | - |
| Year: 2011¹⁴N= 808 Percutaneously inserted CVCsCVCs (334 patients)blood obtained from a peripheral vein, clinical features consistent with bloodstream infection, no other site of infection, and a PICC in place for at least 3complicationsStudy Design: Retrospective cohort studySetting: Level III NICUNon-femoral n = 278 (190 patients)blood obtained from a peripheral vein, clinical features consistent with bloodstream infection, no other site of infection, and a PICC in place for at least 3 days.Catheter-related sepsis Incidence: 134/368 (36.4%)Risk of Bias: LowDates: January 2004 – December 2007Adverse events: Phlebitis: a linear red streak developed along the superficial veins from the insertion site; can be culture negative; patients with both inflammation and phlebitis categorized as phlebitisOld Percutaneous CVC: 8.8 cases per 3 catheter-days• New Percutaneous CVC: 9.9 cases per catheter-days• Premature infants with BW ≤ 1500g | Isai | | · • | | , , , , , , |
| 2011 ¹⁴ Inserted CVCs Inserted CVCs Non-femoral n = 278 (190 Dates: January 2004 – December 2007 Inclusion Criteria: Premature infants with BW ≤ 1500g Dates: January 2004 December 2007 Premature infants with BW ≤ 1500g Dates: January 2004 Catheter-related sepsis Incidence: 134/368 (36.4%) Catheter-related sepsis Incidence: 134/368 (36.4%) Catheter-related sepsis Incidence: 134/368 (36.4%) Catheter-related sepsis Incidence: 134/368 (36.4%) Catheter-related sepsis Incidence: 134/368 (36.4%) Adverse events: | ., | | - | 1 | |
| Study Design: Retrospective cohort study Catheter-related sepsis Incidence: 134/368 (36.4%) Femoral n = 240 (183 infants) Catheter-related sepsis Incidence: 134/368 (36.4%) Old Peripheral CVC: 88/518 (16.9%) Now Peripheral CVC: 46/290 (15.9%) Adverse events: Dates: January 2004 − December 2007 December 2007 Inclusion Criteria: Premature infants with BW ≤ 1500g Non-femoral n = 278 (190 infection, no other site of infection, and a PICC in place for at least 3 days. Adverse events: Phlebitis: a linear red streak developed along the superficial veins from the insertion site; can be culture negative; patients with both inflammation and phlebitis categorized as phlebitis Premature infants with BW ≤ 1500g Non-femoral n = 278 (190 infection, no other site of infection, no other site of infection, and a PICC in place for at least 3 days. Old Peripheral CVC: 46/290 (15.9%) Phelbitis: a linear red streak developed along the superficial veins from the insertion site; can be culture negative; patients with both inflammation and phlebitis categorized as phlebitis New Percutaneous CVC: 8.8 cases per catheter-days New Percutaneous CVC: 9.9 cases per catheter-days New Percutaneous CVC: 9.9 cases per catheter-days New Percutaneous CVC: 9.9 cases per catheter-days Phlebitis categorized as phlebitis Premature infants with an introduction cannula | | • | CVCs (334 patients) | | complications |
| Study Design: Retrospective cohort studySetting: Level III NICUpatients) Femoral n = 240 (183 infants)infection, and a PICC in place for at least 3 days.Incidence: 134/368 (36.4%) • Old Peripheral CVC: 88/518 (16.9%) • New Peripheral CVC: 46/290 (15.9%) • New Peripheral CVC: 46/290 (15.9%) • p = 0.680Risk of Bias: LowDates: January 2004 – December 2007• Old type Percutaneously inserted CVCs used before June 2006— single lumen silicone catheter with an introduction cannulaPhlebitis: a linear red streak developed along the superficial veins from the insertion site; can be culture negative; patients with both inflammation and phlebitis categorized as phlebitis• Old Percutaneous CVC: 8.8 cases per 3 catheter-days • New Percutaneous CVC: 9.9 cases per catheter-days • New Percutaneous CVC: 9.9 cases per catheter-days • p = 0.121 | 2011** | Inserted CVCs | Non formand n 270 (100 | | Cath aton valetad coursis |
| Retrospective cohort study Location: Taiwan Risk of Bias: Low Dates: January 2004 – December 2007 Inclusion Criteria: Premature infants with BW ≤ 1500g Pemoral n = 240 (183 infants) Adverse events: Phlebitis: a linear red streak developed along the superficial veins from the insertion site; can be culture negative; patients with both inflammation and phlebitis categorized as phlebitis Old Peripheral CVC: 88/518 (16.9%) New Peripheral CVC: 46/290 (15.9%) Phlebitis: a linear red streak developed along the superficial veins from the insertion site; can be culture negative; patients with both inflammation and phlebitis categorized as phlebitis New Peripheral CVC: 88/518 (16.9%) Phlebitis: a linear red streak developed along the superficial veins from the insertion site; can be culture negative; patients with both inflammation and phlebitis categorized as phlebitis Premature infants with an introduction cannula | Ctudu Dasian | Setting Lovel III NICH | • | | · · |
| Study Coation: Taiwan Infants Adverse events: | | Setting: Level III NICO | ' | <u> </u> | |
| Risk of Bias: Dates: January 2004 – December 2007 Inclusion Criteria: Premature infants with BW ≤ 1500g Dates: January 2004 – December 2007 • Old type Percutaneously inserted CVCs used before June 2006— single lumen silicone catheter with an introduction cannula • Old type Percutaneously inserted cVCs used along the superficial veins from the insertion site; can be culture negative; patients with both inflammation and phlebitis categorized as phlebitis • p = 0.680 Rate • Old Percutaneous CVC: 8.8 cases per catheter-days • New Percutaneous CVC: 9.9 cases per catheter-days • p = 0.121 | • | Location: Taiwan | ` | uays. | , , , , , , |
| Risk of Bias: Dates: January 2004 – December 2007 • Old type Percutaneously inserted CVCs used before June 2006— single lumen silicone catheter with an BW ≤ 1500g • Old type Percutaneously inserted CVCs used along the superficial veins from the insertion site; can be culture negative; patients with both inflammation and phlebitis categorized as phlebitis • Old Percutaneous CVC: 8.8 cases per catheter-days • Premature infants with BW ≤ 1500g • Premature infants with an introduction cannula • Discourse phlebitis: a linear red streak developed along the superficial veins from the insertion site; can be culture negative; patients with both inflammation and phlebitis categorized as phlebitis • New Percutaneous CVC: 9.9 cases per catheter-days | study | Location: Talwan | illiants) | Adverse events: | , |
| Low December 2007 inserted CVCs used before June 2006— insertion site; can be culture negative; patients with both inflammation and phlebitis categorized as phlebitis December 2007 inserted CVCs used before June 2006— insertion site; can be culture negative; patients with both inflammation and phlebitis categorized as phlebitis catheter-days Old Percutaneous CVC: 8.8 cases per 2006 catheter-days | Rick of Rias: | Dates: January 2004 – | Old type Percutaneously | | 1 |
| before June 2006— Inclusion Criteria: Premature infants with BW ≤ 1500g before June 2006— insertion site; can be culture negative; patients with both inflammation and phlebitis categorized as phlebitis phlebitis categorized as phlebitis insertion site; can be culture negative; patients with both inflammation and phlebitis categorized as phlebitis p = 0.121 | | • | | • | |
| Inclusion Criteria:single lumen siliconepatients with both inflammation andNew Percutaneous CVC: 9.9 cases per• Premature infants with BW ≤ 1500gcatheter with an introduction cannulaphlebitis categorized as phlebitiscatheter-days• p = 0.121 | 2011 | December 2007 | | | • • • |
| Premature infants with BW ≤ 1500g Premature infants with an introduction cannula phlebitis categorized as phlebitis catheter-days p = 0.121 | | Inclusion Criteria: | | | , |
| BW ≤ 1500g introduction cannula • p = 0.121 | | | _ | 1 ' | • • • |
| | | | | Financia dataganzada da princaria | · |
| | | 2 = 20008 | Saasaan sainiala | Thrombosis: leg swelling with or without | ▼ p - 0.121 |
| Fredriction Criteria. | | Exclusion Criteria: | New type n= 290 | | DICC with CPS by Porcutaneous CVC site |
| | | | | Para Para Para Para Para | (recalculated by CDC to show infections per site, |
| (recalculated by edge to show infections p | | • | · | Catheter site inflammation: local site | instead of site infections per all infections) |
| • No PICC needed inflammation with no pathogen identified | | | 2130 200 | | mistead of site infections per all infections) |

| Study Information | Population and Setting | Intervention/ Study Groups | Definitions | Results |
|-------------------|------------------------|--|---|---|
| | Detailed records | Non-femoral n= 120 in 114 | and it was diagnosed in the presence of | • Femoral: 83/410 (20.2%) |
| | unavailable | infants | lymphangitis, purulence, or at least 2 | Non-femoral: 51/398 (21.8%) |
| | | Femoral n = 170 in 111 | signs of inflammation (erythema, | • p = NR |
| | | infants) | tenderness, increased warmth, or | • Adjusted OR for Femoral Placement: 1.53 (1.07 – |
| | | | induration); can be culture negative | 2.25) |
| | | New type | | • p = 0.044 |
| | | Percutaneously inserted | Cholestasis: direct bilirubin level ≥ 1.5 | |
| | | CVCs used since July | mg/dL | PICC with CRS by Percutaneous CVC type |
| | | 2006 due to hospital | | Old Percutaneous CVCs: 88/518 (17.0%) |
| | | policy change – single | Occlusion of the PICC: diagnosis only if it | New Percutaneous CVC: 46/290 (15.9%) |
| | | lumen silicone catheter | happened under standard practice and | • p = NR |
| | | with a stiffening stylet | was excluded if it occurred because of | Adjusted OR for New Percutaneous CVC: 1.18 |
| | | and an Excalibur | misconduct | (0.76 – 1.83) |
| | | introducer | Duratura, computately basis | • p = 0.462 |
| | | | Rupture: completely broken | |
| | | Device/agent: Catheter site | Percutaneous CVC rather than simple | Suspected sepsis |
| | | and catheter type | leakage | Incidence: |
| | | Standard proventive | Extravasation: dislodgement of a PICC | Old Percutaneous CVC: 28/518 (5.4%) |
| | | Standard preventive measures: | Extravasation. dislougement of a Fice | New Percutaneous CVC: 17/290 (5.9%) |
| | | Peripheral CVC usually | Leakage: NR | • p = 0.786 |
| | | placed by a nursing | Pericardial effusion: NR | |
| | | specialist who had | | Topic-specific outcomes: |
| | | worked in this field for | Sampling /Testing strategy: | Duration of indwelling PICC (days): |
| | | more than 15 years. | When clinical symptoms and signs | • Old Percutaneous CVC: 21.0 (11.0-29.0) |
| | | Residents or clinical | developed, a single blood sample | • New Percutaneous CVC: 16.0 (6.75 – 25.0) |
| | | neonatologist fellows | culture was obtained peripherally | • p < 0.001 |
| | | followed a standardized | (never through the Peripheral CVC), | |
| | | insertion procedure | and empiric antibiotic therapy was | Adverse events |
| | | under supervision. | administered. Usually 1 mL (at least | Noninfectious complications |
| | | All Percutaneous CVC | 0.5 mL) of blood was taken for each | Percutaneous CVC without CRS by PICC site |
| | | were inserted through a | culture | • Femoral: 95/410 (23.2%) |
| | | peripheral vein; Tip | | • Non-femoral: 139/398 (34.9%) |
| | | location confirmed to be | Other notes: | • p = NR |
| | | in a central vein | The principle of site selection did not | • Adjusted OR (femoral): 0.76 (0.51–1.15) |
| | | The Percutaneous CVC | change when authors substituted | • p = 0.197 |
| | | were advanced or | new-type Peripheral CVC for the old | Percutaneous CVC without CRS by PICC type |
| | | retreated if needed, | type. | • Old Percutaneous CVC: 135/518 (26.0%) |
| | | after a follow-up chest | In this paper, the authors define PICC | • New Percutaneous CVC: 99/290 (34.1%) |
| | | radiograph was taken. | as percutaneously inserted central | • p = NR |
| | | Standardized procedure | catheter not peripherally inserted | • Adjusted OR (new type): 1.13 (0.74 – 1.71) |
| | | for the insertion and | central catheter. Catheters are | • p = 0.573 |
| | | continuous care of the | inserted into the greater and lesser | Phlebitis |
| | | Percutaneous CVC, | saphenous veins of the lower | • Old Percutaneous CVC: 31/518 (6.0%) |

| Study Information | on Population and Setting Intervention/ Study Groups Definitions Results | | | Results |
|--|---|---|--|--|
| | | regardless of the insertion site. • After successful insertion, 10% povidone-iodine containing alcohol (75%) was applied to the insertion site, normal saline used to decolorize, and the area was covered by a transparent dressing ("Tegaderm"). • Nurses checked the insertion site frequently and changed the dressing every 3 days. • The Percutaneous CVC lines were not impregnated with antibacterial or antiseptic agents and antibiotic lock prophylaxis was not used. | extremities, basilic veins or cephalic veins of the upper extremities, and femoral veins and the tip end in a central vein | New Percutaneous CVC: 9/290 (3.1%) p = 0.072 Thrombosis Old Percutaneous CVC: 0/290 (0%) p = 0.214 Catheter site inflammation Old Percutaneous CVC: 36/518 (6.9%) New Percutaneous CVC: 31/290 (10.7%) p = 0.064 Cholestasis Old Percutaneous CVC: 88/518 (26.3%) New Percutaneous CVC: 50/290 (25.0%) p = 0.739 Occlusion Old PICCs: 37/518 (7.1%) New PICCs: 24/290 (8.3%) p = 0.559 Rupture Old PICCs: 13/518 (2.5%) New PICCs: 13/290 (4.5%) p = 0.127 Extravasation Old PICCs: 8/518 (1.5%) New PICCs: 13/290 (4.5%) p = 0.012 Leakage Old PICCs: 8/518 (1.5%) New PICCs: 8/290 (2.8%) p = 0.235 Pericardial effusion Old PICCs: 0/518 (0%) New PICCs: 1/290 (0.34%) |
| | | | | • p = 0.359 |
| Author: Tsai Year: 2009 ¹³ Study Design Retrospective cohort | Number of patients: N = 334 Number of lines: N= 518 Percutaneously Inserted CVC | Study Groups: Femoral: N = 183 Patients N = 240 Percutaneously Inserted CVCs Non-femoral: N = 190 | Outcome Definitions: Catheter-related sepsis (CRS): culture confirmed; at least 1 positive culture of blood obtained from a peripheral vein, clinical features consistent with bloodstream infection, no other site of | Primary Outcomes: Catheter related sepsis. Incidence • Femoral: 54/240 (22.5%) • Non-femoral: 34/278 (12.2%) • p = 0.002 |
| study Risk of Bias | Setting: Level III NICU | patients N= 278 Percutaneously Inserted CVCs | infection, and a PICC in place for at least 5 days. | Rate • Femoral: 10.9/1000 catheter days |

| Study Information | Population and Setting | Intervention/ Study Groups | Definitions | Results |
|-------------------|--|---|--|--|
| Moderate | Location: Taiwan | | Adverse events: | Non-femoral: 6.8/1000 catheter days |
| | | Device/agent: Catheter type | Phlebitis: a linear red streak developed | • p = 0.012 |
| | Dates: January 2004 – | | along the superficial veins from the | Insertion of PICCs at femoral sites |
| | June 2006 | Standard preventive | insertion site; can be culture negative; | •OR:1.91 (95% CI, 1.17–3.12,) |
| | | measures: | patients with both inflammation and | • p = 0.010) |
| | Inclusion Criteria: | All Percutaneously | phlebitis categorized as phlebitis | |
| | Premature infants with | Inserted CVCs were | Thrombosis: leg swelling with or without | Topic-specific outcomes: |
| | BW < 1500g | single lumen silicone | poor perfusion developed | Duration of indwelling PICC, d (mean ± SD) |
| | | catheters with an | Catheter site inflammation: diagnosed in | • Femoral: 20.7 ± 8.9 |
| | | introduction cannula. | the presence of lymphangitis, purulence, | Non-femoral: 17.0 ± 9.3 |
| | Exclusion Criteria: | Percutaneously Inserted | or at least 2 signs of inflammation | • p < 0.001 |
| | Early death unrelated | CVCs usually placed by a | (erythema, tenderness, increased | |
| | to Percutaneously | nursing specialist who | warmth, or induration); can be culture | Adverse events |
| | Inserted CVCs | had worked in this field | negative | Phlebitis |
| | insertion | for more than 15 years. | Cholestasis: direct bilirubin level ≥ 1.5 | • Femoral: 0/240 (0%) |
| | No Percutaneously | Residents or a clinical | mg/dL | Non-femoral: 29/278 (9.3%) |
| | Inserted CVCs needed | neonatologist fellow | Occlusion of the Percutaneously Inserted | • p < 0.001 |
| | Detailed records | would perform and | CVCs: diagnosis only if it happened under | Thrombosis |
| | unavailable | follow a standardized | standard practice and was excluded if it | • Femoral: 2/240 (0.8%) |
| | | procedure under | occurred because of malpractice | • Non-femoral: 0/278 (0%) |
| | | supervision. | Rupture: completely broken | • p = 0.214 |
| | | Authors used a | Percutaneously Inserted CVCs rather than | Catheter site inflammation |
| | | standardized procedure | simple leakage | • Femoral: 6/240 (2.5%) |
| | | for the insertion and | Extravasation: dislodgement of a | • Non-femoral: 30/278 (13.3%) |
| | | continuous care of the | Percutaneously Inserted CVCs | • p < 0001 |
| | | PICC, regardless of the | Leakage: NR | Cholestasis |
| | | insertion site. | Sampling /Tosting strategy | • Femoral: 49240 (26.7%) |
| | | After successful | Sampling /Testing strategy: | • Non-femoral: 56/278 (29.4%) |
| | | insertion, 10% povidone- | When clinical symptoms and signs developed, a single blood sample | • p = 0.861 |
| | | iodine containing alcohol | culture was obtained peripherally | Occlusion |
| | | (75%) was applied to the | (never through the Percutaneously | • Femoral: 18/240 (7.5%) |
| | | insertion site, normal | Inserted CVCs), and empiric antibiotic | • Non-femoral: 19/278 (6.8%) |
| | | saline used to | therapy was administered. Usually 1 | • p = 0.769 |
| | | decolorize, and the area | mL (at least 0.5 mL) of blood was | Rupture |
| | | was covered by a | taken for each culture | • Femoral: 8/240 (3.3%) |
| | | transparent dressing | taken for each careare | • Non-femoral: 5/278 (1.5%) |
| | | ("Tegaderm"). | Other notes: | • p = 0.265 |
| | | Nurses checked the insertion site frequently | • In this paper, the authors define PICC | Extravasation |
| | | insertion site frequently and changed the | as percutaneously inserted central | • Femoral: 5/240 (2.1%) |
| | | • | catheter not peripherally inserted | • Non-femoral: 3/278 (1.5%) |
| | | dressing every 3 days. • The PICC lines were not | central catheter. Here a | • p = 0.481 |
| | | | Percutaneously Inserted CVCs is a CVC | • ρ = 0.481 Leakage |
| | | impregnated with | in the femoral vein both centrally and | • Femoral: 4/240 (1.7%) |
| | | | and the same and t | • Femoral: 4/240 (1.7%) Page 41 of 137 |

| Study Information Population and Setting Intervention/ Study Groups Definitions Results | | | Results | |
|---|--|---|--|---|
| | | antibacterial or antiseptic agents and antibiotic lock prophylaxis was not used. • The confirmation of catheter-related complications and the decisions for the removal of a PICC, either elective or due to complications were made by the attending neonatologists, or senior residents on duty. | peripherally inserted in inserted catheters where the tip terminated in central veins other than the femoral vein. • Peripheral sites other than femoral veins were preferred over femoral sites. Femoral venous cannulation was performed when all other peripheral vascular accesses failed. • For those with need for early removal, the second PICC line was usually placed at least 3 days after the condition for early removal was resolved. | • Non-femoral: 4/278 (2.3%) • p = 0.555 |
| Author: Hoang | Number of patients: | Study Groups: | Outcome Definitions: | Primary Outcomes: |
| Year: 2008 ²¹ | N = 396 Number of lines: N= 477 PICCs | Upper extremity group: n= 370 PICCs of 183 infants | Catheter related bloodstream infection (CRBSI): [CDC guidelines] positive culture of an intravascular catheter with the | CRBSI: Rate; infections/ 1000 catheter days • UE: 7.1 |
| Study Design | N-477 FICCS | Lower extremity group: | same species as from ≥1 peripheral blood | • LE: 4.8 |
| Retrospective cohort | Setting: Level III NICU | n=107 PICCs of 190 infants | culture. For culture, ≥ 1.0 mL of blood was | • p = NS |
| study | Location: USA | Device/agent: Catheter site | procured from both a peripheral site and | |
| Risk of Bias Low | Dates: June 2002-June 2006 Inclusion Criteria: NR Exclusion Criteria: Neonates with • Liver dysfunction • Inborn errors of metabolism Liver dysfunction: direct hyperbilirubinemia (serum direct bilirubin of >2.0 mg/dL) and high alanine aminotransferase and alanine aminotransferase levels. | Standard preventive measures: Indications for a PICC are determined by the attending neonatologists PICCs are placed by specialized nursing teams supervised by the neonatologists No patient had 2 PICCs at the same time. Heparin routinely added to PICC. | the central lines Adverse events: Mechanical complications were determined whenever dislodgement of a PICC occurred. • Phlebitis: a physicochemical or mechanical complication not related to a proven infection. • Cholestasis & renal insufficiency: elevated direct bilirubin ≥ 2 mg/dL and maximum serum creatinine level of ≥ 1.6 mg/dL, respectively. • Catheter occlusion: pump occlusion or inability to flush and/or withdraw from the PICC and the cause to be related to thrombotic event. • Leakage: construed as fluid extravasation and/or pleural or pericardial effusion. Sampling /Testing strategy: | Incidence, n (%) • UE: 43/370 (11.6%) • LE: 10/107 (9.3%) • p = NS Coagulase-negative Staphylococcus incidence, n (%) • UE: 37/43 (86.0%) • LE: 5/10 (50.0%) • p <0.05 Topic-specific outcomes: Duration of PICC, median (IQR), d • UE: 13.0 (8.0-22.0) • LE: 16.0 (11.0-26.8) • p <0.004 Adverse events: Phlebitis, n (%): • UE: 21/370 (5.7%) • LE: 6/107 (5.6%) • p = NS Cholestasis, n (%): |

| Study Information | Population and Setting | Intervention/ Study Groups | Definitions | Results |
|--------------------------|---------------------------|------------------------------------|---|--|
| | | | For culture, ≥1.0 mL of blood was | • UE: 112/370 (30%) |
| | | | procured from both a peripheral site | • LE: 25/107 (21.5%) |
| | | | and the central lines. | • p < 0.05 |
| | | | | Occlusion, n (%): |
| | | | Other notes: | • UE: 25/370 (6.7%) |
| | | | Lower extremity PICCs were inserted | • LE: 8/107 (7.5%) |
| | | | because of failure to insert PICCs in | • p = NS |
| | | | the upper extremity, or it was the | Leakage, n (%): |
| | | | primary selection site | • UE: 25/370 (6.7%) |
| | | | | • LE: 3/107 (2.8%) |
| | | | | • p = NS |
| | | | | Time to first complication, median (IQR) d: |
| | | | | • UE: 9.0 (4.0–18.0) |
| | | | | • LE: 15.0 (9.5–22.0) |
| | | | | • p = 0.050 |
| Author: Breschan | Number of patients: | Study Groups: | Outcome Definitions: | Primary Outcomes: |
| Addio: bicschan | N= 236 | Internal jugular- group I: | Catheter associated infection (CAI) | Catheter associated infections: |
| Year: 2007 ¹⁷ | Number of lines: | N= 129 internal jugular | diagnosis was made in patients who | Incidence, n (%): |
| rear. 2007 | N = CVCs | venous catheters among | developed signs of infection (fever | • Group I: 20/129 (15.5%); 95% CI: 0.09-0.23 |
| Study Design | N = eves | 103 patients | [<38°C], hypothermia [<36.5°C], | • Group S: 5/107 (4.7%); 95% CI: 0.01-0.11 |
| Retrospective cohort | Setting: NICU | Subclavian- group 2: | leukocytosis or leukopenia, apnea, or | • P < 0.01 |
| study | Jetting. Wes | n=107 subclavian venous | bradycardia) with no other clinically | • Observed RR = 3.29 |
| stady | Location: Austria | catheters among 84 | apparent site of infection. | |
| Risk of Bias | 2000 Maria | neonates | apparent site of infection. | Cox Proportion Hazard Model |
| Low | Dates: 1998- 2006 | coates | Suspected infection: If the tip culture was | Suspected infection: |
| 20.11 | | Device/agent: Catheter site | found to be negative after catheter | Incidence, n (%): |
| | Inclusion Criteria: | | removal, the diagnosis was reversed to | • Group I: 7/129 (5.4%); 95% CI: 0.02-0.12 |
| | Neonates who | Standard preventive | suspected catheter infection | |
| | received a CVC placed | measures: | retrospectively. | • Group S: 4/107 (3.7%); 95% CI: 0.01-0.1 |
| | percutaneously in | Catheter type | , , | • p = 0.38 |
| | either the internal | Standard: 2-French | Adverse events: | Cathotax associated a Suspected infaction. |
| | jugular or the | single-lumen catheter | | Catheter associated + Suspected infection: Incidence, n (%): |
| | subclavian vein while | • Baby > 1.9 kg: 2-French | Clinical obstruction: NR | , , , |
| | undergoing abdominal | single lumen or 4-French | | • Group I: 27/129 (20.9%); 95% CI: 0.14-0.29 |
| | or thoracic noncardiac | double lumen catheter | Clinical thrombosis: NR | • Group S: 9/107 (8.4%); 95% CI: 0.03-0.15 |
| | surgery. | inserted | | • p < 0.01 |
| | Comprised babies who | All CVCs inserted in the | Clinical dislocation: NR | Tonic specific outcomes: |
| | underwent major | operating room during | | Topic-specific outcomes: Length of catheterization in relation to BW: |
| | surgery during their | general anesthesia | Pneumothorax: NR | • Group I: Median: 10 |
| | first 28 days of life or, | before surgery. | | • |
| | if born prematurely, | Insertion was performed | Hemothorax: NR | Group S: Median: 10 Adverse events: |
| | until 28 days had | by three | | Clinical obstruction: |
| | | anesthesiologists | Sampling /Testing strategy: | |
| | | 1 | | • Group I: 8/129 (6.2%); 95% CI: 0.027-0.12 |

| Study Information | Population and Setting | Intervention/ Study Groups | Definitions | Results |
|-------------------|---|--|--|--|
| | elapsed from the | experienced in central | The catheter tips were taken under | • Group S: 1/107 (0.9%); 95% CI: 0.0002-0.05 |
| | calculated birth date. | venous line placement in | sterile conditions to the microbiology | • p < 0.05 |
| | Babies weighing <4.6 | infants. | laboratory where they were plated on | |
| | kg at time of | The vein selected for | 5% horse blood agar. | Clinical thrombosis: |
| | operation. | cannulation was | | • Group I: 1/129 (0.7%); 95% CI: 0.002-0.04 |
| | Availability of | determined by the | Other notes: Infants in Group I (internal | • Group S: 2/107 (1.8%); 95% CI: 0.002-0.06 |
| | patient's tip culture | attending | jugular insertion site) were of younger | • p = 0.43 |
| | after CVC removal. | anesthesiologist. | gestational age and lower birthweight | |
| | | Aseptic technique used | than infants in Group II (subclavian | Clinical dislocation: |
| | Exclusion Criteria: | during all insertions: use | insertion site). Cox Regression analysis for | • Group I: 1/129 (0.7%); 95% CI: 0.0002-0.04 |
| | If percutaneous catheter | of sterile gloves, drapes, | association wit with Catheter-associated | Group S: NR |
| | implantation was | gowns, and facemasks. | infection over time: | • p = 0.54 |
| | unsuccessful in patients | Patient's skin disinfected | • Study group (insertion site): p = 0.002 | |
| | | by rubbing the site of | • Weight: p = 0.075 | Pneumothorax: |
| | | insertion with sterile | Post-conceptual age: p = 0.931 | • Group I: 2 |
| | | gauze soaked in a | | • Group S: 1 |
| | | solution of 2% | | • p = NR |
| | | chlorhexidine in 70% | | Hemothorax: |
| | | alcohol and was allowed | | Group I: 1 |
| | | to dry. | | • Group S: 0 |
| | | Specific catheters were | | • p = NR |
| | | fixed by stitches; No | | |
| | | tunneling was | | |
| | | performed. | | |
| | | Exit site of the CVC | | |
| | | covered by an occlusive | | |
| | | dressing unless the | | |
| | | baby's weight was less | | |
| | | than 1 kg, then | | |
| | | Steristrips were used. | | |
| | | Any manipulations on | | |
| | | the catheters were | | |
| | | performed by NICU | | |
| | | nurses following a | | |
| | | standardized protocol. | | |
| | | Proper catheter tip | | |
| | | positioning in the | | |
| | | superior caval vein was | | |
| | | confirmed by x-ray. | | |
| | | Postoperatively all | | |
| | | babies were cared for in | | |
| | | the (NICU) or | | |
| | | intermediate care unit | | |

| Study Information | Population and Setting | Intervention/ Study Groups | Definitions | Results |
|-----------------------|------------------------|--|--|-----------------------------|
| | , op | for neonates; Both units | | |
| | | were managed by the | | |
| | | same team of doctors | | |
| | | and nurses who had all | | |
| | | been trained in neonatal | | |
| | | intensive care medicine. | | |
| | | Any manipulations on | | |
| | | the catheters were | | |
| | | performed by the NICU | | |
| | | nurses following a | | |
| | | standardized protocol. | | |
| | | Three-way stopcocks | | |
| | | connecting the hub with | | |
| | | the intravenous sets | | |
| | | were changed every 48 | | |
| | | h, or even 24 h when | | |
| | | used for total parenteral | | |
| | | nutrition administration. | | |
| | | Stopcocks and hubs | | |
| | | were disinfected with a | | |
| | | solution of 2% | | |
| | | chlorhexidine in 70% | | |
| | | isopropyl alcohol using a | | |
| | | sterile swab immediately | | |
| | | before and after each | | |
| | | manipulation and | | |
| | | wrapped in sterile gauze | | |
| | | dressing. | | |
| | | Babies weighing less | | |
| | | than 1 kg received a low | | |
| | | dose of vancomycin | | |
| | | prophylactically until the | | |
| | | CVC was in place | | |
| Author: | Number of patients: | Study Groups: | Outcome Definitions: | Primary Outcome: |
| Vegunta ¹⁸ | N = 126 | Neck site group: | Catheter infection NR | Catheter infection: |
| Year: 2005 | Number of lines: | n=88 CVCs implanted in | Line consis/ Cathotox related consis | Incidence, n (%): |
| Study Docion | N = 137 tunneled | NICU | Line sepsis/ Catheter-related sepsis: definition NR | • Neck: 11/88 (12.5%) |
| Study Design | catheters | L/R Subclavian vein | denintion NK | • Groin: 1/49 (2%) |
| Retrospective cohort | Sotting: NICL | L/R Internal jugular vein | Adverse events: | • p = 0.032 |
| study | Setting: NICU | R external jugular vein | | Cath stay valeted sension |
| Risk of Bias | Location: USA | R internal jugular vein | Dislodgement: NR | Catheter-related sepsis: |
| High | LUCATION: USA | Construction at the same | Pleural/pericardial complication: NR | Rate per 1000 catheter days |
| i iigii | | Groin site group: | Fieural/pericardial complication. NK | • Neck: 5.8 |
| | | | | Page 45 of 137 |

| Study Information | Population and Setting | Intervention/ Study Groups | Definitions | Results |
|-------------------|--|---|---|--|
| | Dates: June 1998- | n=49 CVCs implanted in | Clotted catheter: NR | • Groin: 0.7 |
| | February 2003 | NICU | | • p = 0.032 |
| | | L/R Long saphenous vein | Leak from tunnel: NR | |
| | Inclusion Criteria: | | | Topic-specific outcomes: |
| | Infants requiring single | Device/agent: Catheter site | Sampling /Testing strategy: | Catheter live days (mean ± 1 SD) |
| | lumen tunneled | | Line sepsis was confirmed with | Neck: 21.6 (23.8) |
| | catheter during study | Standard preventive | cultures, and salvage was attempted | • Groin: 30.5 (45) |
| | period | measures: | by treating appropriate antibiotics. | • p = 0.105 |
| | | Catheter type | | |
| | Exclusion Criteria: | Single lumen 2.7F | Other notes: | Adverse events: |
| | NR | tunneled catheters used | Infants in the "groin site" group | Total complications (including infections) |
| | | in all neonates | were significantly younger, and of | Incidence, n (%): |
| | | • 3.5F percutaneous | lower birthweight and gestational | • Neck: 26/88 (29.5%) |
| | | introducer sets were | age than infants in the "neck site" | • Groin: 4/49 (8.2%)) |
| | | used for subclavian | group. | • p = 0.005 |
| | | placement. | There were no catheter related deaths | Rate per 1000 catheter days: |
| | | | in this study. | • Neck: 13.7 |
| | | Neck lines mostly | | • Groin: 2.67 |
| | | performed in operating | | • p = 0.005 |
| | | room (OR), placed under | | |
| | | general anesthesia. | | Dislodgement/Accidental removal, n (%): |
| | | Groin lines were | | • Neck: 9/88 (10.2%) |
| | | performed | | • Groin: 0/49 (0%)) |
| | | predominantly in NICU | | • p = 0.050 |
| | | Babies ≥ 1500 g had | | |
| | | attempts at | | Pleural/ pericardial complications, n (%): |
| | | percutaneous subclavian | | • Neck: 4/88 (4.5%) |
| | | access; failing which, | | • Groin: 0/49 (0%)) |
| | | ipsilateral internal or | | Clotted catheter, n (%): |
| | | external jugular vein was | | • Neck: 0/88 (0%) |
| | | accessed by cut down. | | • Groin: 3/49 (6.1%)) |
| | | No patient in this study | | Leak from tunnel, n (%): |
| | | population had 2 tunneled | | • Neck: 2/88 (2.3%) |
| | | catheters concurrently. | | • Groin: 0/49 (0%) |

Page **46** of **137**

Table 21 Risk of Bias of Two Group Studies on Catheter Sites

| Author Year | All study groups derived from similar source/reference populations | Attrition not significantly different across study groups | Measure of exposure is valid | Measure of outcome is valid | Investigator blinded or were outcomes well-defined and objective to endpoint assessment | Potential confounders identified | Statistical adjustment for potential confounders done | Funding source(s) disclosed and no obvious conflict of interest | Overall Risk of Bias |
|---------------------------------|---|---|------------------------------|--------------------------------------|---|--|--|---|----------------------------|
| Bashir 2016 ¹⁹ | ✓ | | ✓ | ✓ | ✓ | ✓ | ✓ | ✓ | Low |
| Breschan 2007 ¹⁷ | ✓ | | ✓ | ✓ | ✓ | ✓ | ✓ | | Low |
| Elmekkawi 2019 ²⁰ | ✓ | ✓ | ✓ | ✓ | ✓ | ✓ | | ✓ | Low |
| Garcia 2019 ¹⁵ | ✓ | ✓ | ✓ | ✓ | ✓ | ✓ | ✓ | ✓ | Low |
| Hoang 2008 ²¹ | ✓ | | ✓ | ✓ | ✓ | ✓ | | ✓ | Low |
| Litz 2017 ¹⁶ | ✓ | ✓ | ✓ | ✓ | ✓ | ✓ | | | Low |
| Tsai 2011 ¹⁴ | ✓ | | ✓ | ✓ | ✓ | ✓ | ✓ | ✓ | Low |
| Tsai 2009 ¹³ | ✓ | | ✓ | ✓ | ✓ | ✓ | | | Moderate |
| Vegunta 2005 ¹⁸ | · | | ✓ | NO | NO | ✓ | | | High |
| Wrightson 2013 ²² | ✓ | | ✓ | ✓ | ✓ | ✓ | | ✓ | Low |

C.4. Number of Catheter Lumens

Key Question 4. In NICU patients requiring umbilical venous catheters, does the use of single-lumen, compared with double-lumen, umbilical venous catheters prevent CLABSI in NICU patients?

Table 22 Summary of Findings on the Number of Umbilical Venous Catheter Lumens to Prevent CLABSI

| Outcome | Findings | Quantity and Type of Evidence and Sample Size | GRADE of Evidence for Outcome and Limitations of the Evidence |
|---------|---|--|---|
| CLABSI* | Two observational studies reported an increase in CLABSI is associated with an increasing number of lumens. One cohort study²³ examining 2,017 UVCs reported an increase in the adjusted risk of CLABSI in patients who had lines with two lumens compared to lines with one lumen (aOR: 2.7 (95% CI: 1.1-6.8); P = 0.04) | 2 OBS n = 4,052 lines ²³ n= 250 lines ¹⁵ | Low |

| Outcome | Findings | Quantity and Type of Evidence and Sample Size | GRADE of Evidence for Outcome and Limitations of the Evidence |
|------------------|--|---|---|
| | One case control study¹⁵ reported a large increase in the adjusted odds of CLABSI in patients with double lumen catheters compared with patients with single lumen catheters, however confidence intervals were wide [OR: 5.8 (95% CI: 1.2 – 30.0); p = 0.03] | | |
| Catheter Sepsis* | One RCT ²⁴ found that no infections were reported in either group. | 1 RCT n=43 lines ²⁴ | Low ● Imprecision: only one study, low number of events |

Table 23 Extracted Information on the Number of Umbilical Venous Catheter Lumens

| Study Information | Population and Setting | Intervention/ Study Groups | Definitions | Results |
|-----------------------------|--|--|---|--|
| Author: Levit ²³ | Number of patients: | Study Groups: | Outcome Definitions: | Primary Outcomes: |
| | N = 2676 patients | UAC: n=2035 | CLABSI: CDC/NHSN definition, and if no | CLABSI: |
| Year: 2020 | Number of lines: | UVC: n=2017 | other source was identified and if the UC | Incidence, n/N (%) |
| | N= 4052 lines | | was still indwelling or had been removed | • UAC: 2/2035 (0.1%) |
| Study | | Double lumen: n=679 | within 48 hours of the onset of infection | • UVC: 19/2017 (0.9%) |
| Design: Cohort | Setting: | Single lumen: n=3373 | | , , , |
| | Level IV NICU | | Adverse events: | UVC: |
| Risk of Bias: Low | | Device/agent: Catheter type; | Complications: break/rupture, | Adjusted incidence rate ratio/ 1000 central-line |
| | Location: USA | Number of lumens | occlusion, catheter tip malposition, poor | days: (adjusted for infant's sex, gestational age, |
| | | | perfusion to lower extremity, CLABSI, | and birthweight) |
| | Dates: January 1, 2008 – May | Standard preventive | thrombus, or effusion | • alRR: 2.7 (95% CI: 1.1-6.8); P = 0.04 |
| | 31, 2018 | measures: | | Adjusted rate/ 100 catheter days |
| | | UC insertion is a sterile, | Sampling /Testing strategy: NR | Double lumen UVC: 2.0 |
| | Inclusion Criteria: | bedside procedure | | Single lumen UVC: 0.7 |
| | Any infant admitted to the | typically performed by | Other notes: Only the first instance of a | |
| | NICU who had a UAC, UVC, | advanced practice | complication within a neonate was | Cumulative incidence of UVC-related CLABSI: |
| | or both successfully placed | providers, pediatric | considered in the analyses. | • First week of life: <1% |
| | (i.e., catheter tip in the | interns and residents, and | | • At day 14: 3.6% |
| | desired, central location) | neonatal-perinatal | | • At day 18: 16.5% |
| | | medicine fellows | | |
| | Exclusion Criteria: | Double-lumen catheter | | Topic-specific outcomes: |
| | • NR | insertion is based solely | | Mean dwell time, days (range) |
| | | on anticipated need | | • UAC: 5.5 days (1-22) |
| | | UVCs used for infusion of | | • UVC: 7.6 days (1-21) |
| | | intravenous fluids, | | • p = NR |
| | | parenteral nutrition and | | - ρ - ττιτ |
| | | lipids and continuous | | Adverse events |
| | | medication infusions; may | | All complications: |
| | | be used for infusion of | | 55 |

Page **48** of **137**

| Study Information | Population and Setting | Intervention/ Study Groups | Definitions | Results |
|------------------------------|---------------------------------|---|---------------------------------------|--|
| | | intermittent medications | | Adjusted incidence rate ratio/ 1000 central-line |
| | | and blood products | | days |
| | | Blood is not typically | | • IRR for any UAC associated complication: 0.3 |
| | | withdrawn from a UVC | | (95% CI: 0.2-0.4) |
| | | UACs used predominantly | | Adjusted UAC complication rate/ 1000 days: |
| | | blood pressure | | • UAC: 4.6 |
| | | monitoring but may be | | • UVC: 17.6 |
| | | used for infusion of | | • p = NR |
| | | intravenous fluids, | | |
| | | parenteral nutrition and | | Incidence, n/N (%) |
| | | lipids | | • UAC: 51/2035 (2.5%) |
| | | Confirmation of UC | | • UVC: 269/2017 (13.3%) |
| 1 | | placement is via | | • p = NR |
| | | thoracoabdominal | | Adjusted rate/ 1000 central-line days |
| | | radiograph | | Double lumen UVC: 17.2 |
| | | Routine, scheduled | | Single lumen UVC: 15.9 |
| | | reconfirmation of UC | | • p = 0.23 |
| | | location is not performed | | |
| | | Heparin at a | | Complications excluding catheter malposition: |
| | | concentration of 1 U ml | | Adjusted rate/ 1000 central-line days |
| | | ¹ of fluid is infused | | • aIRR: 2.3 (95% CI: 1.2-4.6); P = 0.02 |
| | | continuously through all | | Double lumen UVC: 3.8 |
| | | central line lumens | | Single lumen UVC: 1.6 |
| | | Central line tubing utilized | | Adjusted incidence rate ratio/ 1000 central-line |
| | | for parenteral nutrition, | | days |
| | | intralipids, and/or blood | | • IRR: 1.6 (95% CI: 1.02-2.5) |
| | | products is changed every | | Adjusted rate: |
| | | 24 hours | | • UAC: 3.9 |
| | | Tubing utilized only for | | • UVC: 2.4 |
| | | dextrose containing fluids | | • p = NR |
| | | is changed every 96 hours | | |
| | | An assessment of the | | |
| | | continued need for | | |
| | | central access is typically | | |
| A | Name to a street of the street | made at day 5-7 of use | Outron Befficition | Dilusus Outrons |
| Author: Garcia ¹⁵ | Number of patients: | Clase: | Outcome Definitions: | Primary Outcomes: |
| Vac. 2010 | N = 179 patients | CLABSI: n=74 | CLABSI: CDC 2018 definition | Placement site of CVC: |
| Year: 2019 | Number of lines: N=179 lines | Control: | • Patient ≤1 year of age has at least | Internal jugular, n/N (%) |
| Study | IN-T/3 IIII62 | Non-CLABSI: n=105 | one of the following signs or | • OR: 2.7 (95% CI: 1.5-5.1); P = 0.001 |
| Design: Nested case- | Setting: | INUIT-CLADSI. II-103 | symptoms: fever (>38.0°C), | • Case: 43/74 (58.1%) |
| control | Third-care level NICU | Device/agent: Catheter site; | hypothermia (<36.0°C), apnea, or | • Control: 35/105 (33.3%) |
| CONTROL | Timu-care lever ivico | double lumen catheter | bradycardia, and | • p = 0.001 |
| | | double fullieff cathleter | | Subclavian (percutaneous insertion), n/N (%) |

| Dates: 2014 - Inclusi Pat a C sta inc Pat ins CV Cas dia Con nec the | ation: Mexico tes: January | Standard preventive | Organism(s) identified in blood is | • Case: 17/74 (23%) |
|---|---|---------------------|--|--|
| installe | lusion Criteria: Patients with installation of a CVC during their hospital stay at the NICU were included Patients with first CVC installation and those with CVC duration ≥48 hours Cases were neonates diagnosed with CLABSI Controls were those neonates with a CVC during the same period but who did not develop a CLABSI lusion Criteria: ients who had a catheter called in another hospital | measures: NR | (are) not related to an infection at another site, and The same common commensal is identified by a culture or non-culture based microbiologic testing method, from two or more blood specimens collected on separate occasions Adverse events: CLABSI-related mortality: a death directly related to the infection which occurred during active infection event and no other underlying cause of fatal outcome was present Sampling /Testing strategy: Two-set of blood cultures were obtained in patients with a suspected infection Disinfection with 2% iodine-povidone were performed One peripheral blood culture was obtained along with a catheter-drawn blood culture Other notes: None | Control: 27/105 (25.7%) p = 0.67 Saphenous, n/N (%) Case: 7/74 (9.5%) Control: 16/105 (15.2%) p = 0.25 External jugular, n/N (%) Case: 4/74 (5.4%) Control: 7/105 (6.7%) p = 0.98 Upper limb, n/N (%) Case: 1/74 (1.3%) Control: 12/105 (11.4%) p = 0.01 Brachial, n/N (%) Case: 1/74 (1.3%) Control: 5/105 (4.8%) p = 0.21 Lower limb, n/N (%) Case: 1/74 (1.3%) Control: 3/105 (2.8%) p = 0.64 Double-lumen catheter: OR: 10.0 (95% CI: 2.3-44.3); P = 0.0001 Case: 72/74 (97.3%) Control: 82/105 (78.1%) Topic-specific outcomes: CVC indwelling total time >21 days, n/N (%): OR: 2.9 (95% CI: 1.5-5.4); P = 0.001 Case: 37/74 (50.0%) Control: 27/105 (25.7%) Adverse events CLABSI-related mortality, n/N (%) Case: 5/74 (6.8%) Control: NR |
| Author: Khilnani ²⁴ Numb | mber of patients: | Study Groups: | Outcome Definitions: | Primary Outcomes: |

| Study Information | Population and Setting | Intervention/ Study Groups | Definitions | Results |
|-------------------|------------------------------------|----------------------------------|--|--|
| Year: 1991 | Number of lines: | Double lumen umbilical | Catheter related sepsis: two "positive" | Double Lumen: 0/23 |
| | N = 43 | venous catheter: n=23 | blood cultures for the same organism | Single lumen: 0/20 |
| Study Design: | | | obtained at least 24 hours after umbilical | |
| RCT | Setting: Neonatal ICU | Single lumen umbilical | venous catheter insertion. | Topic-specific outcomes: |
| | | venous catheter: n=20 | | Duration of catheterization, mean days (SD): |
| Risk of Bias: | Location: USA | | Sampling /Testing strategy: | Double lumen: 2.9 (±2.0) |
| High | | Device/agent: single or | Catheter tips were also cultured when | Single lumen: 3 (± 1.2) |
| | Dates: NR | double lumen catheter | catheters were removed due to | p = NR |
| | | Monitoring intervention: | suspected catheter-related sepsis. | |
| | Inclusion Criteria: Critically ill | | · | Number of additional IV catheters needed, |
| | neonates requiring an | Standard preventive | Other notes: None | mean catheters (SD): |
| | umbilical venous catheter | measures: | | Double lumen: 0.8 (±0.1) |
| | | A standard umbilical | | Single lumen: 2.3 (± 0.8) |
| | Indications for umbilical | venous catheter insertion | | p<0.05 |
| | venous catheter included | technique was used. | | |
| | hemodynamic instability | Single and double lumen | | Adverse events |
| | resulting from severe birth | 5-Fr radiopaque | | Leak around the catheter site, n (%): |
| | asphyxia, respiratory distress | polyurethane umbilical | | Double lumen: 0/23 (0) |
| | syndrome, sepsis/pneumonia, | venous catheters were | | Single lumen: 1/20 (5) |
| | meconium aspiration | used. | | p = NR |
| | syndrome, or congenital heart | Central venous pressure | | · |
| | disease. | (CVP) was monitored in | | Occlusion of one lumen, n (%): |
| | | patients when the | | Double lumen: 1/23 |
| | Exclusion Criteria: NR | catheter tip was at the | | Single lumen: 0/20 |
| | | inferior vena cava-right | | p = NR |
| | | atrial junction | | · |
| | | Both lumens of the double | | Other mechanical problems: |
| | | lumen umbilical venous | | None observed |
| | | catheters were used at all | | |
| | | times for the infusion of fluids | | Difficulty with catheter insertion: |
| | | and medications. Heparin (0.5 | | None observed |
| | | U/mL) was used in all fluids | | |
| | | infused via the single or the | | |
| | | double lumen umbilical | | |
| | | venous catheters, regardless | | |
| | | of type of fluid infused. | | |

. Page **51** of **137**

Table 24 Risk of Bias for Randomized Controlled Trials on Number of Catheter Lumens

| Author Year | Described as randomized | Randomization appropriately performed | Described as double- blind | Outcome assessor blinded | Study participant blinded | Investigator blinded | Attrition described | Attrition smaller than 10-15% of assigned patients | Attrition appropriately analyzed | Funding source(s) disclosed and no obvious conflict of interest | Overall Risk of Bias |
|--------------------------------|-------------------------------|---|----------------------------------|--------------------------------|---------------------------------|-------------------------|------------------------|---|--|--|-------------------------|
| Khilnani 1991 ²⁴ | ✓ | | | | | | ✓ | * | ✓ | → | High |

Table 25 Risk of Bias for Two Group Studies on Number of Catheter Lumens

| Author Year | Were patients randomly assigned to the study's groups? | For non-randomized trials, did the study employ any other methods to enhance group comparability such as matching, stratification, or statistical methods to adjust for baseline differences? | Did patients in different study groups have similar levels of performance on the outcome of interest and other important factors at the time they were assigned to groups? | Did the study enroll all suitable patients or consecutive suitable patients within a time period? | Was the comparison of interest prospectively planned? | Were the two groups treated/ evaluated concurrently? | Was the study blinded or double- blinded? | Was the funding for this study derived from a source that would not benefit financially from results in a particular direction? | Risk of Bias |
|------------------------------|--|---|--|---|---|--|---|---|-----------------|
| Garcia 2019 ¹⁵ | ✓ | ✓ | ✓ | ✓ | ✓ | ✓ | ✓ | ✓ | Low |
| Levit 2020 ²³ | ✓ | ✓ | ✓ | ✓ | ✓ | ✓ | ✓ | | Low |

C.5. Skin Antisepsis for Catheter Insertion and Maintenance

Key Question 5: In NICU patients requiring skin antisepsis for catheter insertion and maintenance, does alcoholic chlorhexidine, compared with alcoholic povidone-iodine, prevent CLABSI?

Table 26 Summary of Findings on the Use of 2% Alcoholic CHG vs. 10% PI for Catheter Insertion and Maintenance

| Outcome | Findings | Quantity and Type of Evidence (Sample Size) | GRADE of Evidence for Outcome (Limitations of the Evidence) |
|---------|---|---|--|
| CRBSI* | 1 multicenter RCT ²⁵ using 2% CHG in alcohol base vs 10% PI suggested catheter related blood stream infections did not occur in either group. | 1 RCT n= 48 lines ²⁵ | Very Low Indirect: study not conducted in current standard of care, Imprecision: only one study |
| CABSI* | • 1 multicenter RCT ²⁵ using 2% CHG in alcohol base vs 10% PI suggested no difference in catheter associated blood stream infections: 1/24 (4%) vs. 1/24 (4%); p = 0.99. | 1 RCT n= 48 lines ²⁵ | Very Low Indirect: study not conducted in current standard of care Imprecision: only one study |

| Outcome | Findings | Quantity and Type of Evidence (Sample Size) | GRADE of Evidence for Outcome (Limitations of the Evidence) |
|------------------------------------|--|---|--|
| Presumed BSI* | • 1 multicenter RCT ²⁵ using 2% CHG in alcohol base vs 10% PI suggested no difference between BSI rates: 4/24 (17%) vs. 4/24 (17%); p = 0.99. | 1 RCT n= 48 lines ²⁵ | Very low Indirect: study not conducted in current standard of care Imprecision: only one study |
| Septicemia* | • 1 multicenter RCT ²⁵ using 2% CHG in alcohol base vs 10% PI reported septicemia rates to be similar among groups: 7/24 (29%) vs. 9/24 (38%); p = 0.54. | 1 RCT n= 48 lines ²⁵ | Very low • Indirect: study not conducted in current standard of care • Imprecision: only one study |
| Chlorhexidine gluconate absorption | • 1 multicenter RCT ²⁵ reported an increase in CHG absorption following the first and second dressing change for the infants whose absorption level was 13-100 ng mL-1 during catheterization: 6/7 (85.7%). | 1 RCT n= 48 lines ²⁵ | Very ow Indirect: study not conducted in current standard of care Imprecision: only one study |
| Product-related Adverse Events | • 1 multicenter RCT ²⁵ (Garland 2009) using 2% CHG in alcohol base vs 10% PI reported 2% CHG was not associated with an increased risk of contact dermatitis when compared to control group. | 1 RCT n=48 lines ²⁵ | Very low Indirect: study not conducted in current standard of care, Imprecision: only one study |

Table 27 Extracted Information on the Use of Chlorhexidine Skin Antiseptic

| Study Information | Population and Setting | Intervention/ Study Groups | Definitions | Results |
|-----------------------|--|--|--|---------------------------------|
| Author: | Number of patients: | Intervention n= 24 | Outcome Definitions: | Primary Outcomes: |
| Garland ²⁵ | N = 48 | 2% chlorhexidine gluconate | CRBSI: a BSI in which there was | CRBSI, n (%): |
| | Number of lines: | (CHG) in an alcohol-based | concordance between organisms grown | • CHG: 0/24 (0%) |
| Year: 2009 | N = 48 | solution | from the blood and catheter tip | • PI: 0/24 (0%) |
| | | PICC sites cleansed with | | |
| Study Design: RCT | Setting: five Level III NICUs, | ampoules containing 3mL | CABSI: Not defined | Catheter-associated BSI, n (%): |
| | two community hospitals, 3 | of 2% CHG | | • CHG: 1/24 (4%) |
| Risk of Bias: | university teaching hospitals | All peripheral intravenous | BSI without a source: positive peripheral | • PI: 1/24 (4%) |
| Moderate | | catheter sites were | blood culture during time of | • p = 0.99 |
| | Location: USA | cleansed with CHG | catheterization or within 24 h of catheter | · |
| Intervention | | ampules containing 0.67 | removal, clinical signs and symptoms of a | BSI incidence, n (%): |
| Bucket: Skin prep/ | Dates: 2005-2007 | mL of 2% CHG. | BSI, antibiotic therapy for ≥ 7 days and | CHG: 2.8/ 1000 catheter days |
| skin cleansing/ | | | no other documented primary site of | • PI: 3.0/ 1000 catheter days |
| absorption/ CRBSI, | Inclusion Criteria: | Control n=24 | infection | • p = 0.96 |
| BSI, septicemia | Parental informed consent | 10% povidone-iodine (PI) | | ' |
| | Critically ill neonates at | | Presumed BSI: signs and symptoms of | Presumed BSI, n (%): |
| | least 7 days old - <2 | Standard preventive | sepsis with a negative blood culture | • CHG: 4/24 (17%) |
| | months of age who | measures: | Septicemia: Blood culture drawn while | • PI: 4/24 (17%) |
| | required a PICC | Neonates were block | PICC in situ | • p = 0.99 |
| | • Weight > 1500g | randomized to one of two | | |
| | | treatment groups | | Septicemia, n (%): |

| Study Information | Population and Setting | Intervention/ Study Groups | Definitions | Results |
|-------------------|---|--|---|--|
| Study Information | Population and Setting Exclusion Criteria: • ≥ 60 days of age at enrollment • Catheterization ≤ 48 h • Prior discharge home • Conditions of altered skin | Insertion / Study Groups Insertion sites cleansed with appropriate antiseptic before catheter placement Site dressed with polyurethane dressing | Definitions Severe contact dermatitis: dermatitis score of ≥ 2 Absorption: Not defined Sampling /Testing strategy: | Results |
| | Conditions of altered skin integrity | polyuretnane dressing changed weekly while catheter remained in situ. Same antiseptic was used to re-cleansed site with each dressing change All peripheral intravenous catheter sites were cleansed with the same antiseptic used for PICC insertion All catheters were placed using standard sterile techniques with wide barriers Catheter removal decisions made independently by primary care team Catheter sites (PICC and peripheral) inspected daily for the presence and severity of contact dermatitis by a study nurse using a dermatitis severity scale | Dermatitis assessment inspected daily at catheter sites by study nurse using dermatitis severity scale Peripheral blood cultures performed at discretion of primary care team in neonates with signs of sepsis Blood CHG concentrations determined using liquid chromatography with tandem mass spectrometry following catheter placement, just before the first dressing change and immediately after the first dressing change Other notes: Absorption section of study ended early. Only 10 neonates had concentration measured | Adverse Events: Dermatitis: Cutaneous disinfection with 2% CHG was not associated with an increased risk of contact dermatitis when compared to cutaneous scrub with PI. CHG Absorption > 10 ng mL ⁻¹ after 1 st application of antisepsis • 5/10 (50%) 13-100 ng mL ⁻¹ during catheterization • 7/10 (70%) Increased following 1 st and 2 nd dressing change • 6/7 (85.7%) 100 ng mL ⁻¹ after 3 rd dressing change • 1/10 (10%) |

Table 28 Risk of Bias for Randomized Controlled Trials Using Chlorhexidine Skin Antiseptics

| | | | | | | | | | | Funding source(s) | |
|-------------------------------|------------|---------------|------------|----------|-------------|--------------|-----------|-------------------|---------------|-------------------|----------|
| | Described | Randomization | Described | Outcome | Study | | | Attrition smaller | Attrition | disclosed and no | Overall |
| Author | as | appropriately | as double- | assessor | participant | Investigator | Attrition | than 10-15% of | appropriately | obvious conflict | Risk of |
| Year | randomized | performed | blind | blinded | blinded | blinded | described | assigned patients | analyzed | of interest | Bias |
| Garland 2009 ²⁵ | ✓ | √ | NO | | √ | | | | | √ | Moderate |

Page **54** of **137**

C.6. Chlorhexidine Bathing

Key Question 6. In NICU patients requiring central venous catheters, does chlorhexidine bathing, compared with no bathing or bathing with placebo, prevent CLABSI?

Table 29 Summary of Findings on Bathing with 2% CHG Cloths vs. Placebo or No Bathing to Prevent CLABSI

| Outcome | Findings | Quantity and Type of Evidence (Sample Size) | GRADE of Evidence for Outcome (Limitations of the Evidence) |
|-----------------------------------|---|---|--|
| CLABSI* | 1 observational study²⁶ using 2% CHG washcloths for bathing vs no cleansing suggested there was a significant decrease in CLABSI rate per 1000 central line days: 4.28 vs 8.64; Adjusted IRR by weight = 0.49 (95CI: 0.36-0.68); p = 0.0000. 1 observational study²⁷ using 2% CHG-impregnated cloths for routine bathing vs mild soap in NICU patients suggested bathing with CHG-impregnated cloths is associated with a clinically meaningful reduction in CLABSI rates per 1000 CVC days: 2.32 (1.06-4.40) vs 6.17 (4.77-7.85) p = NR (text states NS). Infants > 1000g: 1.28 vs 4.92; Crude IRR= 0.26 (95% CI: 0.07-0.72), p = NR Infants ≤ 1000g, aged ≥28 days: 5.73 vs 8.97; Crude IRR=0.79 (95% CI: 0.15-2.60), p = NR Neonates ≤ 1000g, aged < 28 days: no CHG received during baseline and intervention periods and showed no difference: 8.62 vs 8.57; Crude IRR=1.01 (95% CI: 0.10-5.62); Adjusted IRR by weight = 0.86 (95% CI: 0.17-4.44), p = NR | 2 OBS n= 4,243 patients ²⁶ n=790 patients ²⁷ | Low |
| Lab-confirmed sepsis* | • One observational study ⁵⁶ reported a reduction in the hazard of lab-confirmed sepsis when comparing patients who received a CHG bath with those who did not, however this reduction did not achieve statistical significance in the analysis for either the intervention period [0.48 (95% CI: $0.24 - 0.95$); p = 0.035], but not when analyzing the combined intervention and implementation period [HR: 0.58 (95% CI: $0.31 - 0.11$); p = 0.10] | 1 OBS n = 1,233 patients ⁵⁶ | Very Low • Imprecision: only one study |
| Culture-negative sepsis* | One observational study⁵⁶ reported a reduction in the hazard of culture-negative sepsis when comparing patients who received a CHG bath with those who did not. This reduction did not achieve statistical significance for the intervention period [HR: 1.17 (95% CI: 0.81 – 1.69); p = 0.39] or the combined intervention and implementation period [HR: HR: 1.08 (95% CI: 0.77 – 1.51); p = 0.66] | 1 OBS n = 1,233 patients ⁵⁶ | Very Low • Imprecision: only one study |
| Product-related Adverse Events | 1 observational study²⁶ using 2% CHG washcloths for bathing vs no cleansing reported no local or systemic adverse events. 1 observational study²⁷ using 2% CHG-impregnated cloths for bathing vs mild soap reported no events of dermatitis or adverse events during intervention period. | 2 OBS ^{26, 27} n = 4,243 patients ²⁶ n = 790 patients ²⁷ | Very Low • Imprecision: small number of events |

. Page **55** of **137**

Table 30 Summary of Findings on a Single Bath with 0.25% CHX Cloths vs. Saline Impregnated Cloths vs. No Cleansing to Prevent CLABSI

| Outcome | Findings | Quantity and Type of Evidence (Sample Size) | GRADE of Evidence for Outcome (Limitations of the Evidence) |
|-----------------------------------|--|---|--|
| Culture positive sepsis | • 1 single-center RCT ²⁸ comparing the use of 0.25% free CHX impregnated washcloths vs saline impregnated washcloths or no cleansing suggested there was no difference in the incidence of culture positive sepsis in the first seven days of life among the three groups comparing different agents for use in a single bath: 1/20 (5%) vs. 2/20 (10%) vs. 2/20 (10%); p = 0.53. | 1 RCT N = 60 patients ²⁸ | Low • Imprecision: only one study |
| Clinical sepsis | • 1 single-center RCT ²⁸ comparing the use of 0.25% free CHX impregnated washcloths vs saline impregnated washcloths or no cleansing suggested there was no difference in the incidence of clinical sepsis in the first seven days of life between the three groups: 2/20 (10%) vs. 3/20 (15%) vs 1/20 (5%); p = 0.41. | 1 RCT N = 60 patients ²⁸ | Imprecision: only one study |
| Hypothermia | • 1 single-center RCT ²⁸ comparing the use of 0.25% free CHX impregnated washcloths vs saline impregnated washcloths or no cleansing reported no instances of moderate hypothermia (<36.0°C); and no difference in instances of mild hypothermia/ cold stress (36.0° - 36.4 1°C) at 30 mins: (2/20 (10%) vs 2/20 (10%) vs 0/20 (0%)). | 1 RCT N = 60 patients ²⁸ | Imprecision: only one study |
| Product-related Adverse Events | • 1 single-center RCT ²⁸ of NICU comparing the use of 0.25% free CHX impregnated washcloths vs saline impregnated washcloths vs no cleansing reported none of the infants had skin erythema, fissuring, or crusting. | 1 RCT N = 60 patients ²⁸ | • Imprecision: only one study |

Table 31 Extracted Information on Chlorhexidine Bathing

| Study Information | Population and Setting | Intervention/ Study Groups | Definitions L | Results |
|--------------------------------|---|--|--|--|
| Author: Westling ⁵⁶ | Number of patients: | Study Groups: | Outcome Definitions: | Primary Outcomes: |
| | N = 1,233 | CHG Bathing: n = 864 | Laboratory confirmed sepsis with | Intervention period only |
| Year: 2020 | Number of lines: | Implementation period: n = | pathogen: the day on which a blood | Lab-confirmed Sepsis |
| | N = NR | 28 | culture that grew a pathogenic organism | HR: 0.48 (95% CI: 0.24 – 0.95); p = 0.035 |
| Study Design: | | Intervention period: n = 836 | was drawn, | |
| Prospective Cohort | Setting: NICU | Infants ≥1.5kg who | Culture-negative sepsis: the day on which | Culture-negative Sepsis |
| Risk of Bias: Low | Location: Zambia | received a CHG bath within three days of NICU | a blood culture that did not grow any organism was drawn | HR: 1.17 (95% CI: 0.81 – 1.69); p = 0.39 Death |
| | Dates: NR Inclusion Criteria: | admission, and weekly thereafter. CHG was diluted with sterile water | All-cause mortality prior to NICU discharge Suspected sepsis: the day on which a blood culture was taken (regardless of culture results) | HR: 0.83 (95% CI: 0.56 – 1.23); p = 0.35 Intervention & implementation period only Lab-confirmed Sepsis |
| | Infants ≥1.5 kg infants admitted to the study NICU during the implementation and intervention periods Exclusion Criteria: Infants: | No Bathing: n = 369 Implementation period: n = 170 Intervention period: n = 199 • Infants who did not receive a bath | Laboratory-confirmed sepsis with contaminant organism Sampling /Testing strategy: • Blood cultures Other notes: None | HR: 0.58 (95% CI: 0.31 – 0.11); p = 0.10 Culture-negative Sepsis HR: 1.08 (95% CI: 0.77 – 1.51); p = 0.66 Death HR: 0.94 (95% CI: 0.64 – 1.38); p = 0.75 |

| Study Information | Population and Setting | Intervention/ Study Groups | Definitions L | Results |
|------------------------------|---------------------------------|---|--|---|
| | Born outside the facility | Device: bath with 2% | | Topic-specific outcomes: |
| | From the baseline period | aqueous CHG | | NR |
| | • <1.5 kg. | | | |
| | With suspected sepsis on | Standard preventive | | Adverse events: |
| | the day of admission | measures: | | There were no reports of local or systemic |
| | | • (1) IPC training; | | adverse events due to the use of CHG baths in |
| | | (2) Locally manufactured | | the study period. |
| | | alcohol hand rub; | | |
| | | • (3) Daily IPC reminders | | |
| | | via short messaging | | |
| | | service (SMS); | | |
| | | • (4) Enhanced routine | | |
| | | cleaning of the | | |
| | | environment including | | |
| | | potential reservoirs of | | |
| | | infection (such as sinks | | |
| | | and suction machines) | | |
| | | with a focus on daily | | |
| | | cleaning of high touch | | |
| | | surfaces and moving | | |
| | | | | |
| | | from clean to dirty | | |
| Author: Cleves ²⁶ | Number of patients: | Intervention: n= 1662 new | Outcome Definitions: | Primary Outcomes: |
| | N = 4,243 | central lines inserted | CLABSI: bloodstream infection confirmed | CLABSI incidence, n (%): |
| Year: 2018 | Number of lines: | | by two blood cultures in a patient with a | CHG bath: 65 |
| | N = 4,243 | July 2014- February 2017 | central line in place for > 2 calendar days, | No CHG bath: 75 |
| Study Design: | | July 2014, Chlorhexidine | with ≥1 of the following symptoms: fever | |
| Retrospective, | Setting: Tertiary care hospital | gluconate (CHG) baths | (body temperature >38°C), hypothermia | CLABSI rate / 1000 central line days |
| quasi-experimental | with NICU | implemented in NICU by | (body temperature <36°C), apnea or | • CHG bath: 4.28 |
| study | | Infection Committee | bradycardia. | No CHG bath: 8.64 |
| | Location: Columbia (South | CHG baths performed by | | • Global IRR = 0.49 (95% CI: 0.35-0.70) |
| Risk of Bias: | America) | NICU nurses using 2 | CLABSI ratio: number of central line | Adjusted IRR by weight= 0.49 (95CI: 0.36- |
| Low | 5 | antiseptic body cleansing | infections/ | 0.68) |
| | Dates: January 2012 – | washcloths with 2% CHG | 1000 central line days. | • p = 0.0000 |
| | February 2017 | in a non-alcohol and non- | Dationt days; number of days since high | |
| | Inclusion Criteria: | alkaline base—one cloth | Patient-days: number of days since birth | Handwashing adherence found to be: |
| | • NR | for upper limbs, neck, | Incidence rate ratio (IRR): ND | • Intervention (CHG bath): 86.5% |
| | ▼ INIX | thorax, back and armpits -the other cloth used for | Sampling /Testing strategy: | • Pre-intervention (No CHG bath): 91.8% |
| | Exclusion Criteria: | inferior limbs, gluteus | Blood cultures | |
| | • NR | and groin | - Blood cultures | Topic-specific outcomes: |
| | - 1411 | and groin | Other notes: None | NR |
| | | | | |

| N = 790 Study Design: Retrospective cohort study Setting: Level III NICU in a tertiary care pediatric hospital Low Risk of Bias: Low Dates: April 1, 2009 – March N = 790 144/195 (74%) Safety Network definition CLABSI cases: same as 2009.American National Healthcare Safety Network definition until April 1, 2013, the need for the CVC to have been in place for ≥ 48 hours before CLABSI onset was added to definition Dates: April 1, 2009 – March National Healthcare Safety Network definition until April 1, 2013, the need for the CVC to have been in place for ≥ 48 hours before CLABSI onset was added to definition Dates: April 1, 2009 – March National Healthcare Safety Network definition Until April 1, 2013, the need for the CVC to have been in place for ≥ 48 hours before CLABSI onset was added to definition P = NR (text states NS) | Study Information | Population and Setting | Intervention/ Study Groups | Definitions L | Results |
|--|-----------------------------|--|---|---|---|
| Cleansing on Z ^{rol} day after birth | | | Neonates with BW > | | Adverse events: |
| here the study period. | | | 1000g started daily skin | | There were no reports of local or systemic |
| Nemates with BW C 1000g started biweekly skin cleansing on 7" day after birth | | | cleansing on 2 nd day after | | adverse events due to the use of CHG baths in |
| 1000g started biweekly skin cleansing on 7th day after birth | | | birth | | the study period. |
| skin cleansing on 7th day after birth Control: n=1246 new central lines inserted January 2012 - June 2014 • Skin disinfection performed before insertion of all central lines and for catheter care every seven days or when necessary, with 2% CHG and 70% alcohol solution Standard preventive measures: NR Author: Quach** Number of patients: N=790 Number of patients: N=790 Number of ines: N=790 Number of patients: N=790 Number of patients: N=790 Number of patients: N=790 Number of patients: N=790 Number of ines: N=790 Number of patients: N=790 N=790 Number of patients: N=790 Number of patients: N=790 Number of patients: N=790 Number of pati | | | Neonates with BW < | | |
| skin cleansing on 7th day after birth Control: n=1246 new central lines inserted January 2012 - June 2014 • Skin disinfection performed before insertion of all central lines and for catheter care every seven days or when necessary, with 2% CHG and 70% alcohol solution Standard preventive measures: NR Author: Quach** Number of patients: N=790 Number of patients: N=790 Number of ines: N=790 Number of patients: N=790 Number of patients: N=790 Number of patients: N=790 Number of patients: N=790 Number of ines: N=790 Number of patients: N=790 N=790 Number of patients: N=790 Number of patients: N=790 Number of patients: N=790 Number of pati | | | 1000g started biweekly | | |
| after birth Control: n=1246 new central lines inserted January 2012 - June 2014 • Skin disinfection performed before insertion of all central lines and for catheter care every seven days or when necessary, with 2% CHG and 70% alcohol solution Standard preventive measures: NR Author: Quach²² Year: 2014 Number of patients: N=790 Number of lines: N=790 Study Design: Retrospective cohort study Study Design: Setting: Level III NICU in a tertary care pediatric hospital central cohort study Dates: April 1, 2009 – March 31, 2013 Location: Canada Low Dates: April 1, 2009 – March 31, 2013 Location: Canada Location: Canada Low Dates: April 1, 2009 – March 31, 2013 Location: Canada Location: | | | , | | |
| Ilines inserted January 2012 - June 2014 * Skin disinfection performed before insertion of all central lines and for catheter care every seven days or when necessary, with 2% CHG and 70% alcohol solution Standard preventive measures: NR | | | = - | | |
| Ilines inserted January 2012 - June 2014 * Skin disinfection performed before insertion of all central lines and for catheter care every seven days or when necessary, with 2% CHG and 70% alcohol solution Standard preventive measures: NR | | | | | |
| January 2012 - June 2014 • Skin disinfection performed before insertion of all central lines and for catheter care every seven days or when necessary, with 2% CHG and 70% alcohol solution | | | Control: n=1246 new central | | |
| Author: Quach ⁷⁷ Author: Quach ⁷⁷ Number of patients: N=790 Number of lines: N=790 Number of lines: N=790 Study Groups: N=790 Study Period Study Period Steffing: Level III NICU in a tertiary care pediatric hospital Low Dates: April 1, 2009 – March Intervention Bucket: CHG bathing Inclusion Criteria: He lines in the lines with a CVC admitted to NICU during study period Study period Study period Author: Quach ⁷⁷ N=20 Number of patients: N=790 Number of lines: N=790 Study Groups: N=790 Study April 1, 2012 Setting: Level III NICU in a tertiary care pediatric hospital venous catheter (CVC) and a BW > 1000g bathed with 2% Chlorhexidine gluconate (CHG) impregnated cloth daily Sucket: CHG bathing - All linfants with a CVC admitted to NICU during study period Study Period Study Period - All linfants with a CVC admitted to NICU during study period Study Period Study Period Study Period Study Reversely All lines intravenous catheters that ended at or near the heart or in a great vessel. Number of patient-days: total number of days that patients spent in the NICU Number of CVC-days: total number of days that patients spent in the NICU Number of CVC-days: total number of days of exposure to at least 1 CVC and was collected days and Age) - Study Groups: - Study Groups: - Primary Outcomes: - CLABSI (incidence) - Total CLABSI (incidence) - Total CLABSI rates/ 1000 CVC-days 95% CI) - Baseline (2.106-4.40) - P = NR (text states NS) - Pooled CHG-Bathed infants (separated by BW and Age) - Substitution of Class of exposure to at least 1 CVC and was collected days: - Study Groups: - CLABSI (incidence) - Total CLABSI rates/ 1000 CVC-days 95% CI) - Baseline (2.106-4.40) - P = NR (text states NS) - Pooled CHG-Bathed infants (separated by BW and Age) - Baseline: 6.0 - Intervention: 1.92 | | | lines inserted | | |
| Author: Quach ⁷⁷ Author: Quach ⁷⁷ Number of patients: N=790 Number of lines: N=790 Number of lines: N=790 Study Groups: N=790 Study Period Study Period Steffing: Level III NICU in a tertiary care pediatric hospital Low Dates: April 1, 2009 – March Intervention Bucket: CHG bathing Inclusion Criteria: He lines in the lines with a CVC admitted to NICU during study period Study period Study period Author: Quach ⁷⁷ N=20 Number of patients: N=790 Number of lines: N=790 Study Groups: N=790 Study April 1, 2012 Setting: Level III NICU in a tertiary care pediatric hospital venous catheter (CVC) and a BW > 1000g bathed with 2% Chlorhexidine gluconate (CHG) impregnated cloth daily Sucket: CHG bathing - All linfants with a CVC admitted to NICU during study period Study Period Study Period - All linfants with a CVC admitted to NICU during study period Study Period Study Period Study Period Study Reversely All lines intravenous catheters that ended at or near the heart or in a great vessel. Number of patient-days: total number of days that patients spent in the NICU Number of CVC-days: total number of days that patients spent in the NICU Number of CVC-days: total number of days of exposure to at least 1 CVC and was collected days and Age) - Study Groups: - Study Groups: - Primary Outcomes: - CLABSI (incidence) - Total CLABSI (incidence) - Total CLABSI rates/ 1000 CVC-days 95% CI) - Baseline (2.106-4.40) - P = NR (text states NS) - Pooled CHG-Bathed infants (separated by BW and Age) - Substitution of Class of exposure to at least 1 CVC and was collected days: - Study Groups: - CLABSI (incidence) - Total CLABSI rates/ 1000 CVC-days 95% CI) - Baseline (2.106-4.40) - P = NR (text states NS) - Pooled CHG-Bathed infants (separated by BW and Age) - Baseline: 6.0 - Intervention: 1.92 | | | | | |
| performed before insertion of all central lines and for catheter care every seven days or when necessary, with 2% CHG and 70% alcohol solution Standard preventive measures: NR Author: Quach²² Namber of patients: N=790 Number of lines: N=790 Study Groups: Intervention: n= 195 >35weeks gestation: N=790 Study Design: Retrospective cohort study Evitary care pediatric hospital tertiary care pediatric hospital tertiary care pediatric hospital Low Dates: April 1, 2009 – March 31, 2013 Bucket: OHG bathing Inclusion Criteria: - All infants with a CVC admitted to NICU during study period and dressing change remained unchanged (same as baseline) as well as bathing frequency with the substitution of Class frequency with the | | | January 2012 - June 2014 | | |
| insertion of all central lines and for catheter care every seven days or when necessary, with 2% CHG and 70% alcohol solution Standard preventive measures: NR N=790 Number of patients: N=790 Number of lines: N=790 Number of lines: N=790 Study Design: Retrospective cohort study Charling: Level III NICU in a tertiary care pediatric hospital Low Dates: April 1, 2009 – March linervention Bucket: Hiclusion Criteria: All Infants with a CVC admitted to NICU during study period Inclusion Criteria: All Infants with a CVC admitted to NICU during study period Feducion Criteria: NP New Study Design: Retrospective Cohort study Inclusion Criteria: All Infants with a CVC admitted to NICU during study period Perimary Outcomes: CLABSI (incidence) Total CLABSI (incidence) Total CLABSI (incidence) Total CLABSI (ance) Now Herror Primary Ductomes: CLABSI (incidence) Total CLABSI (ance) Inclusion Criteria: All Infants with a CVC admitted to NICU during study period Inclusion Criteria: All Infants with a CVC admitted to NICU during study period Primary Outcomes: CLABSI (incidence) Total CLABSI (incidence) Total CLABSI (ance) Inclusion Criteria: All Infants with a CVC admitted to NICU during study period Inclusion Criteria: All Infants with a CVC admitted to NICU during study period Primary Diodstream infections: same as 2009 American National Healthcare Safety Network definition CLABSI cases same as 2009-American National Healthcare Safety Network definition CLABSI cases same as 2009-American National Healthcare Safety Network definition CLABSI cases same as 2009-American National Healthcare Safety Network definition CLABSI cases same as 2009-American National Healthcare Safety Network definition CLABSI (ance) Inclusion Criteria: All Infants with a CVC and a BW > 1000g bathed with 2% chlorhexidine gluconate (CHG) impregnated cloth daily Use of CHG for insertion After April 1, 2012 Infants with a cVC and a BW > 1000g bathed with 2% chlorhexidine gluconate (CHG) impregnated cloth daily Use of CHG | | | Skin disinfection | | |
| Initial field of the substitution of Standard preventive measures; NR | | | performed before | | |
| Care every seven days or when necessary, with 2% CHG and 70% alcohol solution | | | insertion of all central | | |
| when necessary, with 2% CHG and 70% alcohol solution Standard preventive measures: NR Author: Quach ²⁷ Number of patients: N=790 Number of lines: N=790 144/195 (74%) Study Design: Retrospective cohort study Risk of Bias: Low Dates: April 1, 2009 – March 1Low Intervention Bucket: CHG bathing When necessary, with 2% CHG for insertion Study Design: Retrospective cohort study Risk of Bias: Low Dates: April 1, 2009 – March 31, 2013 Inclusion Criteria: • All infants with a CVC admitted to NICU during study period Study period when necessary, with 2% CHG for insertion Standard preventive measures: NR Outcome Definitions: Primary bloodstream infections: same as 2009 American National Healthcare Safety Network definition CLABSI (asses: same as 2009 American National Healthcare Safety Network definition until April 1, 2013, the need for the CVC to have been in place for ≥ 48 hours before CLABSI onset was added to definition Gentral lines: intravenous catheters that ended at or near the heart or in a great vessel. All infants with a CVC admitted to NICU during study period Study period With 2% CHG for insertion 144/195 (74%) After April 1, 2012 Infants with central venous catheter (CVC) and a BW > 1000g bathed with 2% chlorhexidine gluconate (CHG) impregnated cloth daily Use of CHG for insertion and dressing change remained unchanged (same as baseline) as well as bathing frequency with the substitution of days that patients spent in the NICU with the substitution of days of exposure to at least 1 CVC and with the substitution of days of exposure to at least 1 CVC and with the substitution of days of exposure to at least 1 CVC and with the substitution of days of exposure to at least 1 CVC and with the substitution of days of exposure to at least 1 CVC and with the substitution of days of exposure to at least 1 CVC and with the substitution of days of exposure to at least 1 CVC and with the substitution of days of exposure to at least 1 CVC and with the substitution of days of exposure to at least 1 CVC | | | lines and for catheter | | |
| Author: Quach²² Author: Quach²² Number of patients: N=790 Year: 2014 Number of lines: N=790 Study Design: Retrospective cohort study Low Location: Canada Low Low Intervention Bucket: CHG bathing CHG and 70% alcohol solution Standard preventive measures: NR Study Groups: Intervention: n= 195 Nymber of lines: N=790 144/195 (74%) Study Design: Retrospective cohort study Low Risk of Bias: Low Location: Canada Low Dates: April 1, 2009 – March 31, 2013 Bucket: CHG bathing CHG bathing CHG and 70% alcohol solution Study Design: N=790 Author: Quach²² Number of patients: N=790 Number of lines: N=790 144/195 (74%) Study Design: CLABSI (incidence) Primary Dutcomes: CLABSI (incidence) Potal = 75 Po Baseline = 46 Intervention: 9 Healthcare Safety Network definition Until April 1, 2013, the need for the CVC Until April 1, 2013, the need for the CVC Until April 1, 2013, the need for the CVC Until April 1, 2013, the need for the CVC Until April 1, 2013, the need for the CVC Adjusted lines: intravenous catheters that ended at or near the heart or in a great vessel. Wimber of patients: All infants with a CVC admitted to NICU during study period Study Period Study Design: CLABSI (incidence) • Total = 75 • Baseline = 46 • Intervention: 9 Total CLABSI rates/ 1000 CVC-days 95% CI) • Baseline (90eld; 6.17 (4.77-7.85) • Intervention: 2.32 (1.06-4.40) • Adjusted IRR = 0.86 (95% CI: 0.63-1.16) • p = NR (text states NS) Pooled CLABSIs rates/ 1000 CVC-days by CHG use (# CLABSIs rates/ 1000 CVC-days) * (# CLABSIs rates/ 1000 CVC-days) • p = NR (text states NS) Pooled CLABSI rates/ 1000 CVC-days) * (# CLABSIs rates/ 1000 CVC-days) • p = NR (text states NS) Pooled CLABSI rates/ 1000 CVC-days) • p = NR (text states NS) Pooled CLABSI rates/ 1000 CVC-days) • p = NR (text states NS) Pooled CLABSI rates/ 1000 CVC-days) • p = NR (text states NS) Pooled CLABSI rates/ 1000 CVC-days) • p = NR (text states NS) Pooled CLABSI rates/ 1000 CVC-days) • p = NR (text states NS) Pooled CLABSI rates/ 1000 CVC-days) • p = NR (text states NS | | | care every seven days or | | |
| Study Design: Study Design: Study Groups: Namber of lines: N = 790 Study Design: Study Design: Study Design: Study Design: Christian BB Solution Study Design: Incrvention Intervention Intervention Intervention Intervention Study Design: Setting: Level III NICU in a cohort study Dates: April 1, 2009 – March Slucket: CHG bathing Study Design: Inclusion Criteria: After April 1, 2012 In | | | when necessary, with 2% | | |
| Author: Quach ²⁷ Number of patients: N=790 Number of lines: N=790 Study Groups: Intervention: n= 195 Study Design: Retrospective cohort study Cohort study Cohort study Cohort study Dates: April 1, 2009 – March Intervention Intervention Intervention Intervention All infants with a CVC admitted to NICU during study period Feduration Criteria: All infants with a CVC admitted to NICU during study period Feduration Criteria: Primary bloodstream infections: same as 2009 American National Healthcare Safety Network definition Capacita (CLABSI cases: same as 2009 American National Healthcare Safety Network definition National Healthcare Safety Network definition Until April 1, 2013, the need for the CVC to have been in place for ≥ 48 hours before CLABSI onset was added to definition Central lines: intravenous catheters that ended at or near the heart or in a great vessel. Inclusion Criteria: • All infants with a CVC admitted to NICU during study period Feduration Criteria: • All infants with a CVC admitted to NICU during study period • Textusion Criteria: • Exclusion Criteria: • Exclusion Criteria: • Exclusion Criteria: • Clarsi (Feduration Stame as bactining infections: same as 2009 American National Healthcare Safety Network definition Until April 1, 2013, the need for the CVC to have been in place for ≥ 48 hours before CLABSI onset was added to definition Central lines: intravenous catheters that ended at or near the heart or in a great vessel. • Dates: April 1, 2009 – March Inclusion Criteria: • All infants with a CVC admitted to NICU during study period • All infants with a CVC admitted to NICU during study period • All infants with a CVC admitted to NICU during study period • Clarsi (Feduration Safety Network definition Central lines: intravenous catheters that ended at or near the heart or in a great vessel. • Number of patient-days: total number of days of exposure to at least 1 CVC and with the substitution of days of exposure to at least 1 CVC and with the properties of the primary Durison | | | CHG and 70% alcohol | | |
| Author: Quach ²⁷ Number of patients: N=790 Year: 2014 Number of lines: N = 790 144/195 (74%) Setting: Level III NICU in a tertiary care pediatric hospital Low Dates: April 1, 2009 – March Bucket: CHG bathing Inclusion Criteria: CHG bathing Measures: NR Study Groups: Intervention: n = 195 > 35 weeks gestation: 2099 American National Healthcare Safety Network definition CLABSI cases: same as 2009.American National Healthcare Safety Network definition until April 1, 2013, the need for the CVC to have been in place for ≥ 48 hours before CLABSI onset was added to definition Central lines: intravenous catheters that ended at or near the heart or in a great vessel. N=700 Number of patients: N=790 Number of plines: N=790 144/195 (74%) Setting: Level III NICU in a tertiary care pediatric hospital Venous catheter (CVC) and a BW > 1000g bathed with 2% chlorhexidine gluconate (CHG) impregnated cloth daily Use of CHG for insertion and dressing change remained unchanged (same as baseline) as well as bathing frequency with the substitution of days that patients spent in the NICU Number of CVC-days: total number of days that patients spent in the NICU Number of CVC-days: total number of days of exposure to at least 1 CVC and with contents the number of days of exposure to at least 1 CVC and with contents the number of days of exposure to at least 1 CVC and with the substitution of days of exposure to at least 1 CVC and with the substitution of days of exposure to at least 1 CVC and with the substitution of days of exposure to at least 1 CVC and with the substitution of days that patients do number of days that patients spent in the NICU Number of CVC-days: total number of days that patients do number of days that patients spent in the NICU on later the content of the cVC on days of exposure to at least 1 CVC and on the content of the cVC on the content of the cVC | | | solution | | |
| Author: Quach ²⁷ Number of patients: N=790 Number of lines: N=790 Number of lines: N=790 Number of lines: N=790 Number of lines: N=790 Study Design: Retrospective cohort study Cohort study Cohort study Cohort study Risk of Bias: Low Dates: April 1, 2009 − March Bucket: CHG bathing Inclusion Criteria: CHG bathing Number of patients: N=790 Number of patients: N=790 Number of lines: N=790 National After April 1, 2012 After April 1, 2012 Infants with central venous catheter (CVC) and a BW > 1000g bathed with 2% chlorhexidine gluconate (CHG) impregnated cloth daily Use of CHG for insertion and dressing change remained unchanged (same as baseline) as bething frequency with the substitution of substitution of CVC-days: total number of days of exposure to at least 1 CVC and was certainly bloodstream infections: same as 2009.American National Healthcare Safety Network definition CLABSI cases: same as 2009.American National Healthcare Safety Network definition until April 1, 2013, the need for the CVC to have been in place for ≥ 48 hours before CLABSI onset was added to definition Central lines: intravenous catheters that ended at or near the heart or in a great vessel. Number of patient-days: total number of days of exposure to at least 1 CVC and was of exposure to at least 1 CVC and was certainly bloodstream infections: same as 2009.American National Healthcare Safety Network definition Until April 1, 2013, the need for the CVC to have been in place for ≥ 48 hours before CLABSI onset was added to definition Central lines: intravenous catheters that ended at or near the heart or in a great vessel. Number of patient-days: total number of days of exposure to at least 1 CVC and was of exposure to at least 1 CVC and with the substitution of days of exposure to at least 1 CVC and was cellified daily and Age) Intervention: 1.92 | | | Standard preventive | | |
| Year: 2014 N=790 Number of lines: N = 790 Study Design: Retrospective cohort study Retriary care pediatric hospital Low Dates: April 1, 2009 – March Bucket: CHG bathing Inclusion Criteria: CHG bathing N=790 National Intervention: n = 195 >35 weeks gestation: 144/195 (74%) After April 1, 2012 Infants with central venous catheter (CVC) and a BW > 1000g bathed with 2% chlorhexidine gluconate (CHG) impregnated cloth daily Use of CHG for insertion and dressing change remained unchanged study period Inclusion Criteria: CHG bathing N=790 National After April 1, 2012 Infants with central venous catheter (CVC) and a BW > 1000g bathed with 2% chlorhexidine gluconate (CHG) impregnated cloth daily Use of CHG for insertion and dressing change remained unchanged (same as baseline) as well as bathing frequency with the substitution of study period Evelusion Criteria: N = 790 After April 1, 2012 Infants with central venous catheter (CVC) and a BW > 1000g bathed with 2% chlorhexidine gluconate (CHG) impregnated cloth daily Use of CHG for insertion and dressing change remained unchanged (same as baseline) as well as bathing frequency with the substitution of CVC-days: total number of days of exposure to at least 1 CVC and space in the NICU and days of exposure to at least 1 CVC and with the substitution of CVC-days: total number of days of exposure to at least 1 CVC and with the substitution of CVC-days of exposure to at least 1 CVC and with the substitution of CVC-days of exposure to at least 1 CVC and with the substitution of CVC-days in the period condition and Healthcare Safety Network definition CLABSI cases: same as 2009. American National Healthcare Safety Network definition LAfter April 1, 2012 Intervention: 9 Total CLABSI (incidence) Intervention: 9 Total CLABSI (incidence) Intervention: 9 Total CLABSI rates/ 1000 CVC-days 95% CI) Entervention: 2.32 (1.06-4.40) | | | measures: NR | | |
| Year: 2014 Number of lines: N = 790 >35sweeks gestation: 144/195 (74%) 2009 American National Healthcare Safety Network definition CLABSI cases: same as 2009.American National 4 Total = 75 Baseline = 46 Intervention: 9 Retrospective cohort study Setting: Level III NICU in a tertiary care pediatric hospital After April 1, 2012 Infants with central venous catheter (CVC) and a BW > 1000g bathed with 2% chlorhexidine gluconate (CHG) impregnated cloth daily use of CHG bathing Location: Canada Healthcare Safety Network definition until April 1, 2013, the need for the CVC to have been in place for ≥ 48 hours before CLABSI onset was added to definition Total CLABSI rates/ 1000 CVC-days 95% CI) Intervention: Dates: April 1, 2009 – March 31, 2013 Study period Setting: Level III NICU in a Infants with central venous catheter (CVC) and a BW > 1000g bathed with 2% chlorhexidine gluconate (CHG) impregnated cloth daily Use of CHG for insertion and dressing change remained unchanged (same as baseline) as well as bathing frequency with the substitution of Central lines: intravenous catheters that ended at or near the heart or in a great vessel. Pooled CLABSI rates/ 1000 CVC-days by CHG use Very Devolution: With 2% chlorhexidine gluconate (CHG) impregnated cloth daily Use of CHG for insertion and dressing change remained unchanged (same as baseline) as well as bathing frequency with the substitution of Number of CVC-days: total number of days that patients | Author: Quach ²⁷ | Number of patients: | Study Groups: | Outcome Definitions: | Primary Outcomes: |
| N = 790 Study Design: Retrospective cohort study Cohort study Cohort study Setting: Level III NICU in a tertiary care pediatric hospital Cohort study Cohort study Cohort study Risk of Bias: Location: Canada Low Dates: April 1, 2009 − March Bucket: CHG bathing Inclusion Criteria: All infants with a CVC admitted to NICU during study period Exclusion Criteria: N = 790 After April 1, 2012 After April 1, 2012 Infants with central venous catheter (CVC) and a BW > 1000g bathed with 2% chlorhexidine before CLABSI onset was added to definition Central lines: intravenous catheters that ended at or near the heart or in a great vessel. Number of patient-days: total number of days of exposure to at least 1 CVC and with the substitution of days of exposure to at least 1 CVC and was collected daily. Safety Network definition CLABSI cases: same as 2009.American National Healthcare Safety Network definition Until April 1, 2013, the need for the CVC and a BW > 1000g bathed with 2% chlorhexidine before CLABSI onset was added to definition Central lines: intravenous catheters that ended at or near the heart or in a great vessel. Number of CVC-days: total number of days of exposure to at least 1 CVC and was collected daily. Exclusion Criteria: NB. Pooled CLABSI rates/ 1000 CVC-days 95% CI) Baseline = 46 Intervention: 9 Total CLABSI rates/ 1000 CVC-days 95% CI) Baseline = 46 Intervention: 9 Total CLABSI rates/ 1000 CVC-days 95% CI) Baseline = 46 Intervention: 9 Total CLABSI rates/ 1000 CVC-days 95% CI) Baseline = 46 Intervention: 9 Total CLABSI rates/ 1000 CVC-days 95% CI) Baseline = 46 Intervention: 9 Total CLABSI rates/ 1000 CVC-days 95% CI) Baseline = 46 Intervention: 9 Total CLABSI rates/ 1000 CVC-days 95% CI) Baseline = 46 Intervention: 9 Total CLABSI rates/ 1000 CVC-days 95% CI) Baseline = 46 Intervention: 2.32 (1.66-4.40) Busseline = 46 Intervention: 9 Total CLABSI rates/ 1000 CVC-days 95% CI) Baseline = 46 Intervention: 9 Baseline = 46 Intervention: 9 Baseline = 46 Intervention | | N=790 | Intervention: n= 195 | Primary bloodstream infections: same as | CLABSI (incidence) |
| Study Design: Retrospective cohort study Setting: Level III NICU in a tertiary care pediatric hospital cohort study Risk of Bias: Low Dates: April 1, 2009 – March Bucket: CHG bathing Inclusion Criteria: • All infants with a CVC admitted to NICU during study period Fixuly in Criteria: • After April 1, 2012 After April 1, 2012 • Infants with central venous catheter (CVC) and a BW > 1000g bathed with 2% chlorhexidine gluconate (CHG) impregnated cloth daily • Use of CHG for insertion and dressing change remained unchanged (same as baseline) as bathing frequency with the substitution of days of exposure to at least 1 CVC and was cellected daily. Fixulsion Criteria: Setting: Level III NICU in a tertiary care pediatric hospital After April 1, 2012 After April 1, 2012 • Infants with central venous catheter (CVC) and a BW > 1000g bathed with 2% chlorhexidine gluconate (CHG) impregnated cloth daily • Use of CHG for insertion and dressing change remained unchanged (same as baseline) as bathing frequency with the substitution of days of exposure to at least 1 CVC and was cellected daily. Exclusion Criteria: Setting: Level III NICU in a tertiary care pediatric hospital After April 1, 2012 After April 1, 2012 After April 1, 2013, the need for the CVC to have been in place for ≥ 48 hours before CLABSI onset was added to definition Central lines: intravenous catheters that ended at or near the heart or in a great vessel. Number of patient-days: total number of days that patients spent in the NICU Number of CVC-days: total number of days of exposure to at least 1 CVC and was cellected daily. Exclusion Criteria: National Healthcare Safety Network definition until April 1, 2013, the need for the CVC to have been in place for ≥ 48 hours before CLABSI rates/ 1000 CVC-days 95% CI) Baseline (pooled): 6.17 (4.77-7.85) End Intervention: 2.32 (1.06-4.40) Adjusted IRR = 0.86 (95% CI: 0.63-1.16) Pooled CLABSI rates/ 1000 CVC-days 95% CI) Baseline (pooled): 6.17 (4.77-7.85) Baseline (pooled): 6.17 (4.77-7.8 | Year: 2014 | Number of lines: | >35weeks gestation: | 2009 American National Healthcare | • Total = 75 |
| Setting: Level III NICU in a tertiary care pediatric hospital cohort study Retrospective cohort study After April 1, 2012 Infants with central venous catheter (CVC) and a BW > 1000g bathed with 2% chlorhexidine gluconate (CHG) impregnated cloth daily Intervention Bucket: CHG bathing Inclusion Criteria: After April 1, 2012 After April 1, 2012 Infants with a CVC admitted to NICU during study period After April 1, 2012 Infants with central venous catheter (CVC) and a BW > 1000g bathed with 2% chlorhexidine gluconate (CHG) impregnated cloth daily Use of CHG for insertion and dressing change remained unchanged (same as baseline) as well as bathing frequency with the substitution of the substitution of the cVC to have been in place for ≥ 48 hours before CLABSI onset was added to definition Central lines: intravenous catheters that ended at or near the heart or in a great vessel. National Healthcare Safety Network definition until April 1, 2013, the need for the CVC to have been in place for ≥ 48 hours before CLABSI onset was added to definition Central lines: intravenous catheters that ended at or near the heart or in a great vessel. Number of patient-days: total number of days that patients spent in the NICU Number of CVC-days: total number of days of exposure to at least 1 CVC and Exclusion Criteria: NP. After April 1, 2012 After April 1, 2012 Infants with central venous catheter (CVC) and a BW > 1000g bathed with 2% chlorhexidine gluconate (CHG) impregnated cloth daily Use of CHG for insertion and dressing change remained unchanged (same as baseline) as well as bathing frequency with the substitution of days of exposure to at least 1 CVC and Exclusion Criteria: NP. After April 1, 2012 Intervention: 2.32 (1.06-4.40) Adjusted IRR = 0.86 (95% CI: 0.63-1.16) Day Clusted IRR = 0.86 (95% CI: 0.63-1.16) Day Clusted IRR = 0.86 (95% CI: 0.63-1.16) After April 1, 2013, the need for the CVC to have been in place for ≥ 48 hours before CLABSI onset was added to definition Central lines: intrave | | N = 790 | 144/195 (74%) | Safety Network definition | Baseline = 46 |
| tertiary care pediatric hospital Infants with central venous catheter (CVC) and a BW > 1000g bathed with 2% chlorhexidine gluconate (CHG) impregnated cloth daily Intervention Bucket: CHG bathing Inclusion Criteria: All infants with a CVC admitted to NICU during study period Evaluation Criteria: CHG bathing Inclusion Criteria: Inclusion | Study Design: | | | CLABSI cases: same as 2009.American | • Intervention: 9 |
| venous catheter (CVC) and a BW > 1000g bathed with 2% chlorhexidine gluconate (CHG) impregnated cloth daily Use of CHG for insertion admitted to NICU during study period venous catheter (CVC) and a BW > 1000g bathed with 2% chlorhexidine gluconate (CHG) impregnated cloth daily Use of CHG for insertion and dressing change remained unchanged (same as baseline) as well as bathing frequency with the substitution of Venous catheter (CVC) and a BW > 1000g bathed with 2% chlorhexidine gluconate (CHG) impregnated cloth daily Use of CHG for insertion and dressing change remained unchanged (same as baseline) as well as bathing frequency with the substitution of Central lines: intravenous catheters that ended at or near the heart or in a great vessel. Number of patient-days: total number of days of exposure to at least 1 CVC and Vers collected daily Pooled CLABSI rates/ 1000 CVC-days by CHG use (# CLABSIs / annual CVC days) Pooled CHG-bathed infants (separated by BW and Age) Baseline (pooled): 6.17 (4.77-7.85) Intervention: 2.32 (1.06-4.40) Adjusted IRR = 0.86 (95% CI: 0.63-1.16) Pooled CLABSI rates/ 1000 CVC-days by CHG use (# CLABSIs / annual CVC days) Number of CVC-days: total number of days of exposure to at least 1 CVC and Vers collected daily Vers of CHG for the search to a least 1 CVC and vers collected daily Vers of CHG for the search to have been in place for ≥ 48 hours before CLABSI onset was added to definition Central lines: intravenous catheters that ended at or near the heart or in a great vessel. Number of patient-days: total number of days of exposure to at least 1 CVC and vers collected daily | Retrospective | Setting: Level III NICU in a | After April 1, 2012 | National | |
| venous catheter (CVC) and a BW > 1000g bathed with 2% chlorhexidine gluconate (CHG) impregnated cloth daily Use of CHG for insertion and dressing change remained unchanged study period Venous catheter (CVC) and a BW > 1000g bathed with 2% chlorhexidine gluconate (CHG) impregnated cloth daily Use of CHG for insertion and dressing change remained unchanged (same as baseline) as well as bathing frequency with the substitution of Central lines: intravenous catheters that ended at or near the heart or in a great vessel. Number of patient-days: total number of days that patients spent in the NICU Number of CVC-days: total number of days of exposure to at least 1 CVC and Versusion Criteria: NR Venous catheter (CVC) and a BW > 1000g bathed with 2% chlorhexidine gluconate (CHG) impregnated cloth daily Use of CHG for insertion and dressing change remained unchanged (same as baseline) as well as bathing frequency with the substitution of Central lines: intravenous catheters that ended at or near the heart or in a great vessel. Number of patient-days: total number of days that patients spent in the NICU Number of CVC-days: total number of days of exposure to at least 1 CVC and Versusion Criteria: NR Pooled CLABSI rates/ 1000 CVC-days by CHG use (# CLABSIs / annual CVC days) Pooled CHG-bathed infants (separated by BW and Age) Baseline (pooled): 6.17 (4.77-7.85) Intervention: 2.32 (1.06-4.40) Adjusted IRR = 0.86 (95% CI: 0.63-1.16) Pooled CLABSI rates/ 1000 CVC-days in the nicular patients spent in the NICU Number of CVC-days: total number of days of exposure to at least 1 CVC and Versusion Criteria: NR Intervention: 2.32 (1.06-4.40) Adjusted IRR = 0.86 (95% CI: 0.63-1.16) Pooled CLABSI rates/ 1000 CVC-days in the nicular patients of the control o | cohort study | tertiary care pediatric hospital | Infants with central | Healthcare Safety Network definition | Total CLABSI rates/ 1000 CVC-days 95% CI) |
| Location: Canada Intervention Dates: April 1, 2009 – March 31, 2013 Inclusion Criteria: All infants with a CVC admitted to NICU during study period Evaluation Criteria: NR Evaluation Criteria: NR Location: Canada and a BW > 1000g bathed with 2% chlorhexidine gluconate (CHG) with 2% chlorhexidine gluconate (CHG) impregnated cloth daily ended at or near the heart or in a great vessel. Number of patient-days: total number of days that patients spent in the NICU Number of CVC-days total number of days of exposure to at least 1 CVC and was callected daily. Evaluation: Canada and a BW > 1000g bathed with 2% chlorhexidine gluconate (CHG) impregnated cloth daily ended at or near the heart or in a great vessel. Number of patient-days: total number of days that patients spent in the NICU Number of CVC-days: total number of days of exposure to at least 1 CVC and was callected daily. Evaluation: Canada • Intervention: 2.32 (1.06-4.40) • Adjusted IRR = 0.86 (95% CI: 0.63-1.16) • p = NR (text states NS) Pooled CLABSIs rates/ 1000 CVC-days of days of exposure to at least 1 CVC and was of exposure to at least 1 CVC and was callected daily. | - | | venous catheter (CVC) | until April 1, 2013, the need for the CVC | |
| before CLABSI onset was added to definition Intervention 31, 2013 Dates: April 1, 2009 – March 31, 2013 Sucket: CHG bathing Use of CHG for insertion and dressing change remained unchanged (same as baseline) as well as bathing frequency study period St | Risk of Bias: | Location: Canada | and a BW > 1000g bathed | to have been in place for ≥ 48 hours | |
| Dates: April 1, 2009 – March 31, 2013 gluconate (CHG) impregnated cloth daily ended at or near the heart or in a great vessel. Number of patient-days: total number of days that patients spent in the NICU as bathing frequency with the substitution of days of exposure to at least 1 CVC and Silver of the part of | Low | | with 2% chlorhexidine | before CLABSI onset was added to | , , , |
| Intervention Bucket: CHG bathing Inclusion Criteria: • All infants with a CVC admitted to NICU during study period • Exclusion Criteria: • CHG bathing Inclusion Criteria: • All infants with a CVC admitted to NICU during study period • Central lines: intravenous catheters that ended at or near the heart or in a great vessel. • Number of patient-days: total number of days that patients spent in the NICU Number of CVC-days: total number of days of exposure to at least 1 CVC and was collected daily. • Exclusion Criteria: NR | | Dates: April 1, 2009 – March | gluconate (CHG) | definition | · · · · · · · · · · · · · · · · · · · |
| Bucket: CHG bathing Inclusion Criteria: • All infants with a CVC admitted to NICU during study period • Use of CHG for insertion and dressing change remained unchanged (same as baseline) as well as bathing frequency with the substitution of days of exposure to at least 1 CVC and • CHG for insertion and dressing change remained unchanged (same as baseline) as well as bathing frequency with the substitution of days of exposure to at least 1 CVC and was collected daily. • Use of CHG for insertion and dressing change remained unchanged (same as baseline) as well as bathing frequency with the substitution of days of exposure to at least 1 CVC and was collected daily. | Intervention | 31, 2013 | impregnated cloth daily | Central lines: intravenous catheters that | p (cont states its) |
| CHG bathing Inclusion Criteria: All infants with a CVC admitted to NICU during study period Inclusion Criteria: All infants with a CVC admitted to NICU during study period Inclusion Criteria: All infants with a CVC admitted to NICU during as bathing frequency with the substitution of days of exposure to at least 1 CVC and was collected daily. Inclusion Criteria: All infants with a CVC admitted to NICU during as bathing frequency with the substitution of days of exposure to at least 1 CVC and was collected daily. Inclusion Criteria: All infants with a CVC admitted to NICU during as bathing frequency with the substitution of days of exposure to at least 1 CVC and least | Bucket: | | | ended at or near the heart or in a great | Pooled CLABSI rates/ 1000 CVC-days by CHG use |
| • All infants with a CVC admitted to NICU during study period • All infants with a CVC admitted to NICU during study period • All infants with a CVC (same as baseline) as well as bathing frequency with the substitution of days of exposure to at least 1 CVC and was collected daily. | CHG bathing | Inclusion Criteria: | and dressing change | vessel. | 1 |
| admitted to NICU during study period (same as baseline) as well as bathing frequency with the substitution of days of exposure to at least 1 CVC and the substitution of days of exposure to at least 1 CVC and on the substitution of days of exposure to at least 1 CVC and the substitution of days of exposure to at least 1 CVC and on the substitution of days of exposure to at least 1 CVC and the substitution of days of exposure to at least 1 CVC and on the substitution of days of exposure to at least 1 CVC and the substitution of the substitution of the substitution of | | All infants with a CVC | 0 0 | Number of patient-days: total number of | |
| study period as bathing frequency with the substitution of days of exposure to at least 1 CVC and Study period as bathing frequency with the substitution of days of exposure to at least 1 CVC and least 1 C | | admitted to NICU during | | | ` ' |
| with the substitution of days of exposure to at least 1 CVC and • Intervention: 1.92 | | study period | , | Number of CVC-days: total number of | <i>5 ,</i> |
| Exclusion Criteria: NP CUC for the pront was collected daily | | | . , | | |
| - Crude Inn. 0.30 (95% Cl. 0.12-0.70) | | Exclusion Criteria: NR | | was collected daily | |
| | | | | | Ci due inn. 0.30 (33% Ci. 0.12-0.70) |

| Study Information | Population and Setting | Intervention/ Study Groups | Definitions L | Results |
|-------------------|------------------------|---|--|---|
| | | • Infants with BW ≤ 1000g | CLABSI rates per 1,000 CVC-days by year: | • Adjusted IRR (for BW): 0.33 (95% CI: 0.15 – |
| | | bathed with mild soap | CLABSI episodes divided by number of | 0.73) |
| | | until day of life 28, then | central line-days times 1,000 | |
| | | 2% CHG-impregnated | Incidence rate ratios (IRRs): compare | BW >1000g, Age=NR (CHG group) |
| | | cloths used (also used as | CLABSIs/1,000 CVC-days during the | Baseline (pooled): 4.92 (36/7323) |
| | | subgroup comparator— | baseline (2009–2012) and intervention | • Intervention: 1.28 (4/3126) |
| | | mild soap used during time not eligible for CHG | (2012–2013) periods | • Crude IRR= 0.26 (95% CI: 0.07-0.72) |
| | | bath) | Sampling /Testing strategy: NR | BW ≤1000g, Age ≥28 days (CHG group) |
| | | Nurses used 2 CHG wipes | | Baseline (pooled): 8.97 (24/2677) |
| | | per infant per bath | Other notes: None | • Intervention: 5.73 (3/524) |
| | | Clinical care protocols | | • Crude IRR: 0.79 (95% CI: 0.15-2.60) |
| | | similar for all infants in | | (00,000 |
| | | the NICU. | | |
| | | | | BW ≤1000g, age <28 days (Non-CHG group) |
| | | Control: n= 595 | | No CHG bathing during baseline and intervention |
| | | Baseline Period: | | periods |
| | | Before April 1, 2012 | | • Baseline (poled): 8.57 (6/700) |
| | | Infants with BW ≤ 1000g | | • Intervention: 8.62 (2/232) |
| | | at gestational age (GA) ≤ | | • Crude IRR= 1.01 (95% CI: 0.10-5.62) |
| | | 28 weeks & chronological | | • Adjusted IRR (for BW) = 0.86 (95% CI: 0.17- |
| | | age (CA) <28 days bathed | | 4.44) |
| | | twice a week with mild | | |
| | | soap and used 2% | | Topic-specific outcomes: |
| | | aqueous CHG for CVC | | NR |
| | | insertion and dressing | | |
| | | change (also used as | | Adverse events: |
| | | subgroup comparator— | | "No dermatitis or adverse events reported |
| | | Not eligible for CHG | | during the 2012-2013 period." |
| | | bath) | | |
| | | Infants with BW ≤ 1000g | | |
| | | at GA ≤ 28 weeks & CA | | |
| | | ≥28 days bathed twice a | | |
| | | week with mild soap and | | |
| | | used 0.5% alcoholic CHG | | |
| | | in 70% alcohol for CVC | | |
| | | insertion and dressing | | |
| | | change | | |
| | | • Infants with BW ≤ 1000g | | |
| | | at GA 29-35 weeks & CA | | |
| | | ≥28 days bathed every | | |
| | | other day with mild soap | | |
| | | and used 0.5% alcoholic | | |
| | | CHG in 70% alcohol for | | Page 50 of 127 |

| Study Information | Population and Setting | Intervention/ Study Groups | Definitions L | Results |
|----------------------|-------------------------------|---|--|----------------------------|
| | | CVC insertion and | | |
| | | dressing change | | |
| | | • Infants with BW > 1000g | | |
| | | at GA 29-35 weeks & CA | | |
| | | of all ages (days) bathed | | |
| | | every other day with mild | | |
| | | soap and used 0.5% | | |
| | | alcoholic CHG in 70% | | |
| | | alcohol for CVC insertion | | |
| | | and dressing change | | |
| | | • Infants with BW > 1000g | | |
| | | at GA >35 weeks & CA of | | |
| | | ages (days) bathed daily | | |
| | | with mild soap and used | | |
| | | 0.5% alcoholic CHG in | | |
| | | 70% alcohol for CVC | | |
| | | insertion and dressing | | |
| | | change | | |
| | | | | |
| | | Standard preventive | | |
| | | measures: | | |
| | | During study period, CHG | | |
| | | used for skin antisepsis | | |
| | | prior CVC insertion and | | |
| | | for dressing change on all | | |
| | | neonates | | |
| Author: | Number of patients: | Intervention: | Outcome Definitions: | Primary Outcomes: |
| Sankar ²⁸ | N = 60 | n= 20 in each | Primary outcome variables were (a) | Culture positive sepsis |
| oua. | Number of lines: | Group A: n=20 | skin condition score at 24 h, days 3 | • CHX: 1/20 (5%) |
| Year: 2009 | N = 60 | cleansing with wipes | and 7 (b) skin temperature at 30 min, | • Saline: 2/20 (10%) |
| . Cu. 1 2003 | 11 00 | containing 0.25% free CHX | 1 and 6 h, and (c) colonization rates of | • No cleansing: 2/20 (10%) |
| Study Design: RCT | Setting: Level III NICU | (.44% CHdG) | the axilla and the groin at 24 and 72 h | • p = 0.53 |
| | Geren in The | (************************************** | after intervention. | p = 0.33 |
| Risk of Bias: | Location: India | Group B: n=20 | Secondary Outcome Definitions | Clinical sepsis |
| Low | 2004.011 maid | Cleansing with wipes | included the incidence of clinical and | • |
| 20 | Dates: August 2005 – February | containing 0% CHX (Saline | culture positive sepsis in the first | • CHX: 2/20 (10%) |
| Intervention | 2006 | cleansing) | week of life. | • Saline: 3/20 (15%) |
| Bucket: bath/ skin | | 0.031131118/ | Culture positive sepsis: infants with | • No cleansing: 1/20 (5%) |
| colonization/ | Inclusion Criteria: | Wipes placed in sealed | symptoms and/or signs suggestive of | • p = 0.41 |
| Sepsis | Preterm infants of 28-36 | plastic packages | sepsis and a positive blood culture | |
| JCP313 | weeks of gestation with | containing 6 of a given | · · · | Topic Specific Outcomes: |
| | birthweights between | type | (with known pathogens and coagulase | |
| | 1001-2000g admitted to | Infants' skin wiped from | negative staphylococcus) | Adverse Events: |
| | ICU/Postnatal ward | - | | Skin condition |
| | ico/Postilatai waru | neck to sole in 5 steps by | | Page 60 |

| • | • Informed written consent | trained staff/resident- 1 | | |
|---|--|---|---|--|
| Inf scc ins ma ski res (cc pre | from 1 parent Exclusion Criteria: Infants with one minute Apgar Icore <4, hemodynamic Instability, congenital Inalformations, generalized Ikin disorder and who needed Icontinuous positive airway Icontinuous positive airway Icontinuous ventilation) | wipe for each step with the 6 th used as a spare Control n=20 Group C: n=20 No skin cleansing Standard preventive measures: Infants randomized within 1-3 hours of age and stratified into two strata based on birth weight: 1501-2000g and 1001 to 1500g Those who carried out the intervention and investigators were blinded All the infants were monitored until the end of the first week of life for features of sepsis Skin condition assessed by observing skin on abdomen and dorsum of hands/feet for drying, erythema, fissuring, scaling etc. using a 9 point grading scale adopted by Darmstadt et al. from Lane et al. | Clinical sepsis: infants with negative cultures but with positive sepsis screen (as per the unit protocol) Cold stress: defined as per standard definitions; Temperature of 36.0-36.4°C Hypothermia: defined as per standard definitions. Sampling /Testing strategy: Clinical thermometer measured skin temperature—kept in the axilla for 3 min. Other notes: None | None of the infants had skin erythema/ fissuring/ crusting. Median skin condition scores of the three groups were identical at 24, 72, and 168 hours after intervention. Skin temperature: Axillary temperature (°C) Mean skin temperature (sd) Baseline CHX: 36.6 (0.13) Saline: 36.6 (0.13) No cleansing: 36.6 (0.16) p = 0.78 30 mins CHX: 36.6 (0.20) Saline: 36.6 (0.12) No cleansing: 36.7 (0.24) p = 0.46 1 hour CHX: 36.6 (0.13) Saline: 36.6 (0.08) No cleansing: 36.7 (0.14) p = 0.46 6 hours CHX: 36.7 (0.12) Saline: 36.7 (0.07) No cleansing: 36.7 (0.11) p = 0.66 Incidences of hypothermia No instances of hypothermia (<36°) in any group. Incidence of cold stress No infant had cold stress at 1 and 6 hours. 30 mins CHX: 2/20 (10%) Saline: 2/20 (10%) No cleansing: 0 (0%) p = 0.34 Adverse Events: NR |

Page **61** of **137**

Table 32 Risk of Bias of Randomized Controlled Trials on Chlorhexidine Bathing

| | | | | | | | | | | Funding source(s) | |
|------------------------------|------------|---------------|------------|----------|-------------|--------------|-----------|-------------------|---------------|---------------------|---------|
| | Described | Randomization | Described | Outcome | Study | | | Attrition smaller | Attrition | disclosed and no | Overall |
| Author | as | appropriately | as double- | assessor | participant | Investigator | Attrition | than 10-15% of | appropriately | obvious conflict of | Risk of |
| Year | randomized | performed | blind | blinded | blinded | blinded | described | assigned patients | analyzed | interest | Bias |
| Sankar 2009 ²⁸ | ✓ | ✓ | ✓ | ✓ | ✓ | ✓ | | | | | Low |

Table 33 Risk of Bias of Two Group Studies on Chlorhexidine Bathing

| Author Year | All study groups derived from similar source/ reference populations | Attrition not significantly different across study groups | Measure of exposure is valid | Measure of outcome is valid | Investigator blinded to endpoint assessment or outcomes are objective | Potential confounders identified | Statistical adjustment for potential confounders done | Funding source(s) disclosed and no obvious conflict of interest | Overall Risk of Bias |
|--------------------------------|--|---|------------------------------|-----------------------------|---|--|--|---|----------------------------|
| Cleves 2018 ²⁶ | ✓ | ✓ | ✓ | ✓ | | ✓ | | ✓ | Low |
| Quach 2014 ²⁷ | ✓ | ✓ | ✓ | ✓ | | ✓ | ✓ | ✓ | Low |
| Westling 2020 ⁵⁶ | ✓ | ✓ | ✓ | ✓ | | | | ✓ | Low |

C.7. Catheter Hub Manipulation

Key Question 7: In NICU patients with central line catheters does minimizing the number of times central line hubs are accessed prevent CLABSI?

Table 34 Summary of Findings on Catheter Manipulation to Prevent CLABSI in NICU Patients

| Outcome | Findings | Quantity and Type of Evidence (Sample Size) | GRADE of Evidence for Outcome (Limitations of the Evidence) |
|---|--|---|--|
| Catheter-associated bloodstream infection | • 1 single-center observational study ²⁹ reported catheter hub manipulations that required disinfection, disconnection, or drawing blood through central line were associated with an increased risk of infection (OR: 1.2; 95% CI: 1.1 – 1.3). | | Imprecision: Only one study |

Table 35 Extracted Information on Catheter Manipulation

| Study Information | Population and Setting | Intervention/ Study Groups | Definitions | Results |
|-------------------------------|---------------------------|----------------------------|----------------------|--------------------------------------|
| Authors: Mahieu ²⁹ | Number of patients: N=223 | C: n=357 Catheters | Outcome Definitions: | Primary Outcomes: |
| | Number of lines: | | | CABSI incidence per catheter, n (%): |

| Study Information | Population and Setting | Intervention/ Study Groups | Definitions | Results |
|--------------------|----------------------------------|--|--|---|
| Year: 2001 | N=357 | Device/agent: NA | Catheter associated bloodstream | • CABSI: 17/357 (4.8%) |
| | | | infection (CABSI): | • No CABSI: 340/357 (95.2%) |
| Study Design: | Setting: Neonatal ICU | Monitoring intervention: NA | 1) Primary bloodstream | • p = NR |
| Prospective cohort | | | infection according to the CDC | · |
| study | Location: Belgium | Standard preventive | surveillance definition: | Topic-specific outcomes: |
| | | measures: | a) recognized pathogen isolated from | Catheter duration, mean days (SD): |
| Risk of Bias: | Dates: November 1, 1993- | Aseptic technique: An | blood culture | • CABSI: 20.1 (17.5) |
| Low | October 31, 1994 | aseptic technique was | or a skin contaminant isolated from | • No CABSI 9.2 (6.8) |
| | | used during insertion and | two blood cultures drawn on separate | • p < 0.001 |
| | Inclusion Criteria: All neonates | repositioning; this | occasions, | · |
| | with one or more central | included surgical | b) one of following | Disinfection of catheter exit-site, mean no. of |
| | venous catheters admitted to | scrubbing with 4% | clinical signs of infection (fever >38°C, | catheter manipulations (SD): |
| | the NICU. | chlorhexidine, sterile | hypothermia <37°C, apnea or | • CABSI: 5.5 (13.2) |
| | | gloves, drapes, gowns, | bradycardia) and | • No CABSI 12.6 (13.3) |
| | Exclusion Criteria: NR | and facemasks. | | • p < 0.001 |
| | | | 2) Central venous catheter present at the | · |
| | | Skin cleaning: Before | time the blood culture was obtained. | Disinfection of catheter hub, mean no. of |
| | | inserting a catheter, the | | catheter manipulations (SD): |
| | | skin was cleaned with a | Catheter manipulations were stratified | • CABSI: 18.2 (16.2) |
| | | solution of 2% | according to the type of manipulation: | • No CABSI: 7.6 (7.0) |
| | | chlorhexidine in 70% | (1) Disinfection (catheter hub and/or exit | • p < 0.001 |
| | | isopropyl alcohol. | site), (2) connection of an infusion line to the | |
| | | | catheter (glucose solution, parenteral | Administration of glucose solutions, mean no. |
| | | The exit-site of non- | nutrition solution, continuous | of catheter manipulations (SD): |
| | | umbilical central venous catheters was covered | intravenous (IV) medication | • CABSI: 4.7 (6.3) |
| | | | (3) administration of IV drugs in shot | • No CABSI: 2.7 (3.1) |
| | | with a sterile gauze help | (heparin, antibiotics, other), | • p = 0.14 |
| | | in place by an occlusive transparent dressing. | (4) transfusions (plasma, packed red | |
| | | transparent dressing. | blood cells, platelets), | Administration of parenteral nutrition, mean |
| | | The exit-site of umbilical | (5) manipulation of the calibrated fluid | no. of catheter manipulations (SD): |
| | | lines remained uncovered | chamber (addition of electrolytes, | • CABSI: 12.2 (16.1) |
| | | and was cleaned thrice | hypertonic glucose) and finally, | • No CABS: 4.3 (6.7) |
| | | daily with a solution of 2% | (6) blood drawings through the central | • p < 0.05 (=0.02) |
| | | chlorhexidine in 70% | line | |
| | | isopropyl alcohol prior to | | Administration of continuous IV drugs, mean |
| | | the application of a | Adverse events: NR | no. of catheter manipulations (SD): |
| | | powder containing | | • CABSI: 7.1 (6.4) |
| | | virginiamycin. | Sampling /Testing strategy: | • No CABSI: 2.8 (5.7) |
| | |] , , , , , , , , , , , , , , , , , , , | Swabs were taken from the catheter exit | • p < 0.05 (<0.001) |
| | | Line maintenance: Three- | site and hub at the time of sepsis | |
| | | way stopcocks connecting | evaluation as well at catheter removal in | Administration of antibiotics, mean no. of |
| | | the hub with the IV sets | those catheters not associated with | catheter manipulations (SD): |
| | | changed every 48 hours | infection. | • CABSI: 11.6 (17.6) |

| Study Information | Population and Setting | Intervention/ Study Groups | Definitions | Results |
|-------------------|------------------------|---|--|---|
| | | or every 24 hours when | | • No CABSI: 4.6 (8.2) |
| | | used for TPN | A culture was taken from the skin | • p = 0.05 |
| | | administration. The | catheter junction with another sterile | |
| | | stopcocks and hubs were | cotton swab after removal of the | Administration of heparin solution, mean no. of |
| | | disinfected with a | dressing. | catheter manipulations (SD): |
| | | homemade solution 2% | | • CABSI: 7.8 (15.1) |
| | | chlorhexidine in 70% | Other notes: None | • No CABSI: 3.1 (6.4) |
| | | isopropyl alcohol using a | | • p = 0.10 |
| | | sterile swab immediately | | |
| | | before and after each | | Administration of other IV drugs as bolus, mean |
| | | manipulation and | | no. of catheter manipulations (SD): |
| | | wrapped in sterile gauze | | • CABSI: 10.7 (16.8) |
| | | dressing. | | • No CABSI: 3.9 (6.9) |
| | | | | • p = 0.11 |
| | | Gloves and masks were | | |
| | | not used during catheter | | Transfusions, mean no. of catheter |
| | | manipulation, but hands | | manipulations (SD): |
| | | were disinfected with 70% | | • CABSI: 0 (0) |
| | | isopropyl alcohol before | | • No CABSI: 0.4 (3.9) |
| | | and after each catheter manipulation. | | p = "No association" |
| | | Catheters were flushed | | |
| | | with heparinized saline | | Fluid chamber manipulation, mean no. of |
| | | daily at the tie of IV set | | catheter manipulations (SD): |
| | | change. In arterial lines, a | | • CABSI: 0.6 (1.1) |
| | | continuous infusion of a | | • No CABSI: 0.8 (1.9) |
| | | heparinized solution was | | • p = "No association" |
| | | used to control patency. | | |
| | | Antibiotics: not used | | Blood drawing of blood gases, mean no. of |
| | | prophylactically but only | | catheter manipulations (SD): |
| | | for treatment of | | • CABSI: 12.8 (23.5) |
| | | suspected infections. | | • No CABSI: 5.0 (11.9) |
| | | · | | • p < 0.05 (= 0.02) |
| | | Administration of blood | | |
| | | products: No blood | | Blood drawing of others, mean no. of catheter |
| | | products were | | manipulations (SD): |
| | | administered through the | | • CABSI: 3.2 (5.3) |
| | | CVC | | • No CABSI: 1.3 (2.9) |
| | | | | • p < 0.05 (= 0.02) |
| | | | | Number of manipulations, mean no. (SD): |
| | | | | • CABSI: 70.7/100.4 (70.4) |
| | | | | • No CABSI: 28.7/107.9 (26.6) |
| I | | | | • p < 0.001 |
| | | | | |

| Study Information | Population and Setting | Intervention/ Study Groups | Definitions | Results |
|-------------------|------------------------|----------------------------|-------------|---|
| | | | | Manipulation-related risk factors significantly |
| | | | | associated with CLABSI: Multivariable analysis |
| | | | | Disinfection of the catheter hub: |
| | | | | OR: 1.2 (95% CI: 1.1-1.3); SE: 0; p = 0.002 |
| | | | | Blood sampling/drawing (other than blood |
| | | | | gases): |
| | | | | OR: 1.4 (95% CI: 1.1-1.8); SE: 0; p = 0.009 |
| | | | | 1-7 blood samples: |
| | | | | OR: 1.04 (95% CI: 0.33-3.27); p = 0.95 |
| | | | | 8-14 blood samples: |
| | | | | OR: 5.82 (95% CI: 1.53-22.63); p = 0.006 |
| | | | | >14 blood samples: |
| | | | | OR: 8.4 (95% CI: 0-67.1); p = 0.036 |
| | | | | Risk of CLABSI increased with number of blood |
| | | | | samples taken through the central line |
| | | | | Heparinization: |
| | | | | OR: 0.9 (95% CI: 0.8-1.0); SE: 0; p = 0.047 |
| | | | | Antisepsis of exit-site: |
| | | | | OR: 0.9 (95% CI: 0.8-1.0); SE: 0; p = 0.014 |
| | | | | Adverse events: NR |

Table 36 Risk of Bias for Two Group Studies on Catheter Hub Manipulation

| | All study groups | | | | Investigator blinded or | | | | |
|------------------------------|------------------|------------------|-------------|------------|-------------------------|-------------|------------------|---------------------|---------|
| | derived from | Attrition not | Measure | Measure | were outcomes well- | | Statistical | Funding source(s) | |
| | similar | significantly | of | of | defined and objective | Potential | adjustment for | disclosed and no | Overall |
| Author | source/reference | different across | exposure is | outcome is | to endpoint | confounders | potential | obvious conflict of | Risk of |
| Year | populations | study groups | valid | valid | assessment | identified | confounders done | interest | Bias |
| Mahieu 2001 ²⁹ | ✓ | ✓ | ✓ | ✓ | ✓ | ✓ | ✓ | | Low |

C.8. Central Line Antimicrobial Locks

Key Question 8: In NICU patients with central line catheters, does the use of central line antimicrobial locks, compared with standard of care, prevent CLABSI?

Page **65** of **137**

Table 37 Summary of Findings on Antimicrobial Locks vs. Standard of Care to Prevent CLABSI

| Outcome | Findings | Quantity and Type of Evidence | GRADE of Evidence for Outcome and Limitations of the Evidence |
|--|--|--|---|
| Catheter –related bloodstream infection* | Three RCTs found the use of antimicrobial lock prophylaxis was associated with a reduced risk for CR-BSI. Each study used a different antibiotic agent and a different lock protocol. One study³⁰ found the use of Amikacin-heparin locks for 20 minutes two times a day was associated with reduced risk for definite CR-BSI. OR: 0.27 (95% CI: 0.16 – 0.87); p<0.001 One study³¹ found the use of Fucidic acid-heparin locks once per day for 30-60 minutes was associated with reduced risk for CR-BSI. RR: 0.09 (95% CI: 0.01 – 0.72); p<0.01 One study³² found the use of Vancomycin-heparin locks for 20 minutes in neonates who were being fed primarily by parenteral hyperalimentation and for 60 minutes when enteral feeding exceeded 20 mL/kg/day was associated with reduced risk for CR-BSI OR: 0.05 (95% CI: 0.003 – 0.95); p = 0.05* | 3 RCT n=103 ³¹ n=85 ³² n=83 ³⁰ | Moderate • Indirectness: studies not conducted in current standard of care |
| Suspected/ probable bloodstream infection | Three studies reported no difference in suspected or probable CR-BSI with any type of antimicrobial catheter lock | 3 RCT n=103 ³¹ n=85 ³² n=83 ³⁰ | Moderate Indirectness: studies not conducted in current standard of care |
| Hypoglycemia | One study³² reported an increase in hypoglycemia with use of heparin saline infusions (p = 0.03) Two studies^{30, 31} reported that antimicrobial catheter locks were not associated with increased risk for hypoglycemia | 3 RCT n=103 ³¹ n=85 ³² n=83 ³⁰ | Moderate • Indirectness: studies not conducted in current standard of care |
| Antimicrobial resistance | Two studies reported no incidences of resistance to the antimicrobial used in the lock protocol were detected. | 2 RCT n=85 ³² n=83 ³⁰ | Low Indirectness: studies not conducted in current standard of care Imprecision: low number of events |

Table 38 Extracted Information on Central Line Antimicrobial Locks

| Study Information | Population and Setting | Intervention/ Study Groups | Definitions | Results |
|----------------------|---------------------------------|-----------------------------|--|---|
| Author: | Number of patients: | Intervention group B: n=41 | Outcome Definitions: | Primary Outcome: |
| Seliem ³⁰ | N=83 | Amikacin-heparinized saline | <u>Definite Catheter related bloodstream</u> | Definite catheter-related bloodstream |
| | Number of lines: | lock for 20 minutes 2x/ day | infection: When a positive peripheral | infection, n (%): |
| Year: 2010 | N = 83 | | blood culture (through venous puncture) | Amikacin Lock 3/41 (7.3%) |
| | | Control group A: n=42 | concomitant with positive blood culture | • No Lock: 11/42 (26.2%) |
| Study design: RCT | Setting: Level III Neonatal ICU | Heparinized-normal saline | withdrawn from the catheter or catheter | • RR: 0.27 (95% CI: 0.16 – 0.87); |
| | | lock for 20 minutes 2x/ day | tip cultures grew the same species in the | • p < 0.001 |
| | Location: Egypt | | presence of clinical manifestations of | |
| Risk of bias: Low | | Device/agent: Amikacin | sepsis without apparent source of | Probable catheter-related bloodstream |
| | Dates: February 2007- | | bloodstream infection rather than UVC. | infection, n (%): |
| | February 2008 | Monitoring intervention: NR | | • Amikacin Lock 1/41 (2.4% |

| Study Information | Population and Setting | Intervention/ Study Groups | Definitions | Results |
|-------------------|--|--|---|---|
| Study Information | Inclusion Criteria: All neonates (term and preterm) admitted to the unit and were expected to require a UVC for at least 48 hours. Exclusion Criteria: Neonates with indwelling UVCs for more than 24 hours without a lock technique and those who have received systemic antibiotic therapy or were transferred to other hospitals in the first day of life. | Standard preventive measures: Maximum sterile barriers including use of sterile gloves, gown, cap, mask, and a large sterile drape. The umbilical stump and surrounding skin area of at least 5 cm radius were disinfected with 10% povidone iodine prior to catheter insertion. The umbilical stump was cleansed routinely on a daily basis with 70% alcohol. The intravenous tubing was changed every 24 hours using strict sterile technique. Catheter hubs were cleansed with 70% alcohol whenever hubs were accessed. Catheters removed whenever their use was deemed unnecessary. | Probable CR-BSI: Considered when the positive peripheral blood culture and positive blood culture withdrawn from the catheter grew different species. If there were positive cultures from the blood withdrawn from the catheter or catheter tip while peripheral blood culture was sterile in presence of clinical manifestations of infection. Bloodstream infection (BSI) without a source: Positive peripheral blood culture with clinical manifestations of sepsis and negative blood culture withdrawn from the catheter or tip culture. Hypoglycemia: defined as a bedside whole-blood glucose concentration <45 mg/dL Sampling /Testing strategy: All study subjects had a culture taken after 48 hours for early detection of catheter contamination and on the 5th and 10th days. When the UVC was removed, the catheter hubs and distal 5 cm of each catheter were cultured semiquantitatively. Surveillance rectal and axillary cultures were obtained at study entry and at the time of catheter removal. If sepsis was suspected, two blood cultures were obtained (peripheral and central) and a culture from the catheter hub was performed. Susceptibility of bacterial isolates to amikacin was tested for growth on amikacin-containing agar. Evidence of growth indicated resistance. For amikacin group only: serum concentrations of amikacin were measured with fluorescence polarization immunoassay | Results No Lock: 1/42 (2.3%) RR: 1.01 (95% CI: 0.8 − 1.1); p = 0.9 Total Definite and probable catheter-related bloodstream infection, n (%): Amikacin Lock 4/41 (9.7%) No Lock: 12/42 (28.5%) RR: 0.34 (95% CI: 0.02 − 0.65); p = 0.01 BSI without a source, n (%): Amikacin Lock 2/41 (4.9%) No Lock (saline heparin): 2/42 (4.8%) RR: 1.02 (95% CI: 0.76 − 1.12); p = 0.97 All BSI, n (%): Amikacin Lock 6/41 (14.6%) No Lock (saline heparin): 14/42 33.3%) RR: Relative Risk: 0.43 (95% CI: 0.12 − 0.61); p = 0.02 Topic-specific outcomes: Duration of catheter, days, mean (SD) Amikacin Lock 11.6 (2.1) No Lock (saline heparin):10.3 (3.6) Standardized Mean Difference: -0.44 (95% CI: -0.880.004) p = 0.048* Adverse events Mortality, n (%): Amikacin Lock 4/41 (9.8%) No Lock (saline heparin): 8/42 (19.0%) Hypoglycemic episodes, n (%): Amikacin Lock 5/41 (12.2%) No Lock (saline heparin): 8/42 (19.0%) p = 0.27 Portal or IVC thrombosis: None observed |

| Study Information | Population and Setting | Intervention/ Study Groups | Definitions | Results |
|-------------------------------|---|--|---|--|
| | | | Other notes: None | Amikacin resistance: None of the positive cultures grew microorganisms resistant to amikacin and there were no amikacin-resistant microorganisms detected in any skin or rectal surveillance cultures in either group. |
| Author: Filippi ³¹ | Number of patients: N = 103 | Study Groups | Outcome Definitions: | Primary Outcomes: |
| | Number of lines: N = 103 | Intervention group A: N=50 | Definite catheter related bloodstream | Definite catheter-related bloodstream infection |
| Year: 2007 | | Fusidic acid-heparin lock for | infection: considered as one positive | Fusidic acid lock: 1/50 (2%) |
| Study design: RCT | Setting: Neonatal ICU | 30–60 mins, once per day | blood culture in a neonate with CVC, with concordant colonization of catheter hub | Heparin saline: 11/53 (20.8%) Relative Risk: 0.09 (95% CI: 0.01 – 0.72); |
| | Location: Italy | Control group C: n=53 | or tip, clinical manifestations of infection, | • p < 0.01 |
| Risk of bias: | | Heparin-normal saline lock | and no other apparent source for | |
| Moderate I | Dates: July 2004 – Nov. 2005 | for 30–60 mins, once per day | bloodstream infection except CVC. | <u>Suspected catheter-related bloodstream</u> infection |
| ı | Inclusion Criteria: All admitted | Device/agent: Fusidic acid | Suspected CR-BSI: positive culture of | • Fusidic acid lock: 2/50 (4%) |
| | neonates who required a | | catheter hub or tip, clinical | • Heparin saline: 2/53 (3.8%) |
| | nonmedicated CVC for ≥24 hrs. | Monitoring intervention: NA | manifestations of infection, and no other | • Relative Risk: 1.06 (95% CI: 0.16 – 7.24); |
| | | a | apparent source for bloodstream | • p = NS |
| | Exclusion Criteria: Neonates | Standard preventive | infection except CVC, with negative or | |
| | with medicated CVCs and neonates who had CVCs | measures: Catheters were inserted with sterile | not concordant blood culture. | Total Catheter-related bloodstream infection |
| | removed within 24 hrs. or | technique. Skin surface | blood culture. | rate/ 1000 catheter days |
| | were transferred to other | surrounding the insertion | Colonization: positive culture of catheter | Fusidic acid lock: 6.6 |
| | hospitals or died in the first | point was disinfected with | hub or tip with neither concordant blood | Heparin saline: 24.9 |
| | day of life. | 10% povidone-iodine. | culture nor clinical | Relative Risk: 0.28 (95% CI: 0.13 – 0.60); |
| | , | | signs of infection. | • p < 0.01 |
| | | A transparent polyurethane | Non catheter related sepsis: positive | |
| | | dressing was used to cover | blood culture with clinical manifestations | Colonization |
| | | the insertion site. Intravenous | of infection but negative culture of | • Fusidic acid lock: 3/50 (6%) |
| | | tubing was changed daily, and | catheter hub or tip. | • Heparin saline: 2/53 (4%) |
| | | catheter hubs were cleansed | | • Relative Risk: 1.59 (95% CI: 0.28 – 9.12); |
| | | with 2% chlorhexidine every | Hypoglycemia: >180 or <40 mg/dL | • p = NS |
| | | time they were accessed. | Sampling /Testing strategy: In both | Non-catheter-related bloodstream infection |
| | | | groups, cultures of aspired fluid were | • Fusidic acid lock: 4/50 (8%) |
| | | | taken every 2 days before lock | Heparin saline: 4/53 (7.5%) |
| | | | administration for early detection of | • Relative Risk: 1.06 (95% CI: 0.28 – 4.01); |
| | | | catheter contamination. If any clinical | • p = NS |
| | | | sign of CR-BSI was present, two blood | Topic-specific outcomes: |
| | | | cultures were obtained (1 ml specimen | Total catheter days |
| | | | from peripheral vein, 0.5 ml specimen | • Fusidic acid lock: 456 |
| | | | from the catheter) and a culture was | Heparin saline: 522 |
| | | | performed from the catheter hub. In case | • p = NS |
| | | | | |

| Study Information | Population and Setting | Intervention/ Study Groups | Definitions | Results |
|-------------------------------|----------------------------------|------------------------------|---|---|
| • | | | the CVC was removed, hubs and tip (3-4 | Adverse events |
| | | | cm, distal part) were cultured. | Mortality |
| | | | Other notes: None | Fusidic acid lock: 13/50 (26%) (0 with CR-BSI) Heparin saline: 11/53 (20.75%) (4 with CR-BSI) |
| | | | | Treatment-related adverse events: None observed Phototherapy, n Fusidic acid lock: 34/50 (68%) Heparin saline: 35/53 (66.0%) |
| | | | | • Relative Risk: 1.03 (95% CI: 0.77 - 1.38) |
| | | | | Phototherapy, days, mean (±SD) • Fusidic acid lock: 3.1±1.9 • Heparin saline: 2.6±1.3 |
| | | | | <u>Jaundice</u> • Fusidic acid lock: 33/50 (66%) • Heparin saline: 33/53 (62.3%) • Relative Risk: 1.03 (95% CI: 0.77 - 1.38) |
| | | | | Leukopenia: No cases observed |
| | | | | Thrombocytopenia: No cases observed |
| | | | | Sideroblastic anemia: No cases observed |
| | | | | Hypoglycemia: No cases observed |
| Author: Garland ³² | Number of patients: | Study Groups: | Outcome Definitions: | Infections: |
| | N = 85 | Intervention group: n=42 | Definite Catheter related bloodstream | Definite catheter-related bloodstream |
| Year: 2006 | Number of lines: | Vancomycin-heparin saline | infection: a positive peripheral blood | infection, n(%): |
| | N = 85 | lock solution for 20 minutes | culture with concordant colonization of | Vancomycin lock: 0/42 |
| Study design: RCT | | in neonates who were being | the catheter hub or catheter tip. | • Heparin saline: 8/43 (18.6%) |
| | Setting: Level III Neonatal ICU | fed primarily by parenteral | | • Relative Risk: 0.41 (95% CI: 0.08 – 2.00); p = |
| Risk of bias: Low | | hyperalimentation and for 60 | Probable CR-BSI: Defined | 0.006 |
| | Location: USA | minutes when enteral feeding | by either (1) a positive peripheral blood | • OR: 0.05 (95% CI: 0.003 – 0.95); |
| | Dates: May 2000- May 2001 | exceeded 20 mL/kg/day | culture for coagulase negative staphylococci, with concordant | • p = 0.05* |
| | Inclusion Criteria: All neonates | Control group: n=43 Heparin | colonization of the catheter hub or hub | Probable catheter-related bloodstream |
| | who were admitted to the unit | normal saline lock solution | tip, but DNA subtyping was not done or | infection, n (%): |
| | and would require a catheter | for 20 minutes in neonates | (2) a blood culture through the catheter | • Vancomycin lock: 2/42 (4.8%) |
| | | who were being fed | was positive (peripheral culture sterile or | • Heparin saline: 5/43 (11.6%) |

| Study Information | Population and Setting | Intervention/ Study Groups | Definitions | Results |
|-------------------|----------------------------------|--------------------------------|--|---|
| | (newly placed PICC) for at least | primarily by parenteral | not done) for the same organism | • Relative Risk: 0.41 (95% CI: 0.08 – 2.00); |
| | 48 hours. | hyperalimentation and for 60 | recovered from the catheter hub or tip, | • p = 0.43 |
| | | minutes | with clonal concordance confirmed by | |
| | Exclusion Criteria: NR | when enteral feeding | DNA subtyping when the blood culture | Catheter-related bloodstream infection rate/ |
| | | exceeded 20 mL/kg/day | grew coagulase-negative staphylococci | 1000 catheter days |
| | | | | Vancomycin lock: 2.3 |
| | | Device/agent: NR | Bloodstream infection (BSI) without a | Heparin saline: 17.8 |
| | | | source: Defined by a positive peripheral | • Relative Risk: 0.13 (95% CI: 0.01 – 0.57); |
| | | Monitoring intervention: NR | or line blood culture and no other | • p = 0.004 |
| | | | identifiable primary site of infection. | ' |
| | | Standard preventive | Neonates were treated with at least 7 | BSI without a source, n (%): |
| | | measures: Catheters | days of systemic antibiotic therapy. | • Vancomycin lock: 5/42 (11.9%) |
| | | were inserted percutaneously | Cultures of the catheter were negative | • Heparin saline: 5/43 (11.6%) |
| | | by staff neonatologists using | or, when positive, showed colonization | • Relative Risk: 1.02 (0.32-3.28); |
| | | maximal sterile barriers, | with a strain or strains different from | • p = 0.97 |
| | | including a sterile mask, cap, | those recovered from the blood culture. | p 6.57 |
| | | gloves and | | Topic-specific outcomes: NR |
| | | gown, and a large sterile | Adverse events | Adverse events |
| | | drape. Insertion sites were | Hypoglycemia: defined as a bedside | Patients with organ systems affected: None |
| | | disinfected | whole-blood glucose concentration <40 | observed |
| | | with 10% povidone-iodine, | mg/dL | |
| | | and catheters | | Hypoglycemia, n (%): |
| | | were dressed with a | Sampling /Testing strategy: Surveillance | • Vancomycin lock: 8/42 (19.0%) |
| | | polyurethane film dressing. | rectal and axillary cultures were obtained | • Heparin saline: 18/43 (41.9%) |
| | | | at study entry and at time of catheter | • p = 0.03 |
| | | Catheter sites were cleansed | removal. Gram-positive bacterial isolates | p 5.05 |
| | | and redressed on a weekly | that were recovered from catheter | Colonization by vancomycin-resistant gram |
| | | basis or as needed if the | insertion sites, catheter cultures, or | positive bacteria: None observed |
| | | dressing became loose or the | blood cultures were also tested for | |
| | | site wet. Intravenous tubing | resistance to vancomycin. | Minimum inhibitory concentration of gram |
| | | was changed every 3 days | Microorganisms that showed | positive isolates from skin, catheter or blood >2 |
| | | when used for | growth on vancomycin-containing agar | ug/mL: None observed |
| | | hyperalimentation and every | were considered resistant. | |
| | | 24 hours when used for | | Detectable blood vancomycin level >2 μg/mL |
| | | intralipid therapy. Needless | When infants showed signs suspicious for | • Vancomycin lock: 1/42 (2.4%) |
| | | access ports were not used | sepsis, blood cultures were obtained: a 1- | Heparin saline: 0/43 |
| | | during the trial. Catheter | mL specimen drawn by percutaneous | , , , , , , |
| | | hubs were cleansed with | venipuncture and at least 0.5 mL drawn | |
| | | alcohol whenever the hub | through the infant's catheter; the | |
| | | was accessed. | catheter hub was also cultured, using a | |
| | | | premoistened sterile cotton swab. | |
| | | | Catheters were removed at the | |
| | | | discretion of the attending neonatologist. | |
| | | | At that time, a 1-cm x | |

| Study Information | Population and Setting | Intervention/ Study Groups | Definitions | Results |
|-------------------|------------------------|----------------------------|--|---------|
| | | | 1-cm area of skin surrounding the catheter, the catheter hub, and the distal 5 cm of the catheter each were cultured semi quantitatively. Other notes: None | |

Table 39 Risk of Bias for Randomized Controlled Trials on Central Line Antimicrobial Locks

| Author Year | Described as randomized | Randomization appropriately performed | Described as double-blind | Outcome assessor blinded | Study participant blinded | Investigator blinded | Attrition described | Attrition smaller than 10-15% of assigned patients | Attrition appropriately analyzed | Funding source(s) disclosed and no obvious conflict of interest | Overall Risk of Bias |
|-------------------------------|-------------------------|---|---------------------------|--------------------------------|---------------------------------|-------------------------|---------------------|--|--|---|----------------------------|
| Seliem 2010 ³⁰ | ✓ | ✓ | | | ✓ | | ✓ | | | ✓ | Moderate |
| Filippi 2007 ³¹ | ✓ | | | | | | ✓ | | | | High |
| Garland 2005 ³² | ✓ | | √ | | √ | ✓ | √ | √ | | √ | Low |

C.9. Optimal Umbilical Arterial and Venous Catheter Dwell Time

Key Question 9 In NICU patients requiring an umbilical catheter, what is the optimal duration of umbilical artery and umbilical venous catheters to prevent CLABSI?

Table 40 Summary of Findings on the Optimal Duration of Umbilical Catheters Prior to Prevent CLABSI

| Outcome | Findings | Quantity and Type of Evidence and Sample Size | GRADE of Evidence for Outcome and Limitations of the Evidence |
|---------|--|--|---|
| CLABSI* | Three observational studies^{2, 23, 33} found that longer use of umbilical catheter was associated with an increased risk for CLABSI, at seven days of life. One observational study³³ found an increase in the odds of developing a CLABSI for UVCs in situ >7 days (OR: 5.48 (95% CI: 1.18-25.50); p = 0.03). One observational study³⁴ implemented a QI initiative directing providers to increase the dwell time of UVCs from the average of 5 days to 7 days prior to inserting a PICC and found no increase in UVC-associated CLABSI (IRR 1.13 (95% confidence interval 0.469–2.332) P = 0.92) with a 37.5% reduction in replacement with PICCs. One observational study²³ suggested the cumulative incidence of CLABSI increases with increasing UVC dwell time. Cumulative incidence was <1% in the first week of life, 3.6% at day 14, and 16.5% at day 18. | 4 OBS n=986 lines ³³ n=6,000 lines ² n=201 lines ³⁴ n=4,052 lines ²³ | Low |

| Outcome | Findings | Quantity and Type of Evidence and Sample Size | GRADE of Evidence for Outcome and Limitations of the Evidence |
|-----------------------------|---|--|---|
| | One observational study ² suggested CLABSI rates increased beyond 4 days (UVC: 116/2668 (4.3%) vs PICC: 287/3332 (8.6%) p<0.01). For UVCs that were removed, there was more than five times the risk of CLABSI on days 6-7 than on days 4-5. However, this was not reported as statistically significant. UVCs replaced with PICCs before 4 days were associated with a trend of reduced CLABSI in the first PICC, compared with UVCs replaced on or after 4 days. After adjusting for gestational age, this trend continued but no longer reached statistical significance. | | |
| Catheter-related infection* | • One RCT study ³⁵ found the use of umbilical catheter for up to 28 days was associated with higher rate of infections when compared with UVC dwell times of 7-10 days, but the difference was not statistically significant (OR: 1.66; 95% CI: 0.79 – 3.48; p = 0.18). | 1 RCT n=210 lines ³⁵ | Moderate • Imprecision: only one study |
| Sepsis* | One observational study¹² found the incidence of sepsis was higher in umbilical artery catheters in situ for ≥8 days when compared with those in situ for ≤7 days. (13.6% vs. 1.3%; p<0.0001). This study noted an increase in the incidence of sepsis in UVCs in situ for 4-7 days when compared with those in situ for 1-3 days but the UVC numbers were insufficient for valid statistical analysis. | 1 OBS n=2,316 lines ¹² | Very Low ● Imprecision: only one study |
| Adverse Events | One RCT study³⁵ found there was no difference in adverse events between UVCs left in situ for up to 28 days compared with UVCs in situ for 7-10 days. Adverse events included thrombosis, mortality, arrhythmia, embolus, hemorrhage, and pleural effusion One observational study²³ reported a decrease in the rate of adverse events for UVCs compared with UVCs [IRR: 0.3 (95% CI: 0.2-0.4)] | 1 RCT n=210 lines ³⁵ 1 OBS n = 4,052 lines ²³ | Moderate • Inconsistency |

Table 41 Summary of Findings on the Optimal Duration of Umbilical Artery Catheter for Removal to Prevent CLABSI

| | | Quantity and Type of Evidence | GRADE of Evidence for Outcome |
|----------------|---|--|---|
| Outcome | Findings | and Sample Size | and Limitations of the Evidence |
| CLABSI* | • One observational study ²³ reported two CLABSI for 2,035 UAC lines. No conclusions can be drawn about the impact of duration on CLABSI risk. | 1 OBS n = 4,052 lines ²³ | Very Low • Imprecision: only one study |
| Sepsis* | One observational study¹² found the incidence of sepsis was higher in umbilical artery catheters in situ for ≥8 days when compared with those in situ for ≤7 days. (13.6% vs. 1.3%; p<0.0001). | 1 OBS ¹² n=1,699 lines | Very Low ● Imprecision: only one study |
| Adverse Events | • One observational study ²³ reviewed data on 2,035 UAC lines and reported an increase in adverse events with increasing dwell time for UACs. The incidence of complications was 2.5% by day 5, 4.3% by day 10, and 37% by day 20. The most common adverse events were breakage/ rupture (20%), occlusion (10%), and catheter tip malposition (10%). | 1 OBS n = 4,052 lines ²³ | Very Low ● Imprecision: only one study |

Table 42 Summary of Findings on the Optimal Duration Prior to Removal of Umbilical Venous Catheters to Prevent CLABSI

| | | Quantity and Type of Evidence | GRADE of Evidence for Outcome |
|---------|--|--|---|
| Outcome | Findings | and Sample Size | and Limitations of the Evidence |
| CLABSI* | One observational study² suggested CLABSI rates increased beyond 4 days (UVC: 116/2668 (4.3%) vs PICC: 287/ 3332 (8.6%) p<0.01). For UVCs that were removed, there was more than five times the risk of CLABSI on days 6-7 than on days 4-5. However, this was not reported as statistically significant. One observational study³⁴ implemented a QI directing providers to increase the dwell time of UVCs from the average of 5 days to 7 days prior to inserting a PICC and found no increase in UVC-associated CLABSI (IRR 1.13 (95% confidence interval 0.469–2.332;) P = 0.92) with a 37.5% reduction in replacement with PICCs. | 2 OBS n = 1,392 lines ² n = 201 lines ³⁴ | Very Low Consistency: Inconsistent results across studies Imprecision: only one study, low number of events |
| Sepsis* | One observational study¹² found an increase in the incidence of sepsis in UVCs in situ for 4-7 days when compared with those in situ for 1-3 days but the UVC numbers were insufficient for valid statistical analysis (p<0.0001). | 1 OBS n = 2,316 lines ¹² | Very Low • Imprecision: only one study, low number of events |

Table 43 Summary of Findings on the Optimal Duration Umbilical Venous Catheter for Replacement with a Long-term Catheter to Prevent CLABSI

| Outcome | Findings | Quantity and Type of Evidence and Sample Size | GRADE of Evidence for Outcome and Limitations of the Evidence |
|-----------------------------|--|---|---|
| CLABSI* | Two observational studies^{2, 33} found that longer use of umbilical catheter prior to replacement with a PICC was associated with an increased risk for CLABSI. One observational study³³ found an increase in the odds of developing a CLABSI for UVCs in situ >7 days (OR: 5.48 (95% CI: 1.18-25.50); p = 0.03). One observational study² found that the HR of CLABSI increased beyond 3-4 days of dwell time, and the risk doubled every 2 days thereafter if the UVC was followed by PICC insertion (UVC: 116/2668 (4.3%) vs PICC: 287/ 3332 (8.6%) p<0.01). One observational study³⁴ implemented a QI directing providers to increase the dwell time of UVCs from the average of 5 days to 7 days prior to inserting a PICC and found no increase in UVC-associated CLABSI (IRR 1.13 (95% CI: 0.469–2.332); P = 0.92) with a 37.5% reduction in replacement with PICCs. | 3 OBS n = 986 lines ³³ n = 6,000 lines ² n = 201 lines ³⁴ | Low |
| Catheter-related infection* | • One RCT study ³⁵ found the use of umbilical catheter for up to 28 days was associated with higher rate of infections when compared with UVC in place for 7-10 days prior to replacement with a PICC, but the difference was not statistically significant (OR: 1.66 (95% CI: 0.79 – 3.48); p = 0.18). | n = 210 lines ³⁵ | Moderate • Imprecision: only one study |
| Adverse Events | One RCT study³⁵ found there was no difference in adverse events between UVCs left in situ for up to 28 days compared with UVCs in situ for 7-10 days. Adverse events included thrombosis, mortality, arrhythmia, embolus, hemorrhage, and pleural effusion. | 1 RCT n = 210 lines ³⁵ | Moderate • Imprecision: only one study |

Page **73** of **137**

Table 44 Extracted Information on Umbilical Catheter Dwell Time

| Study | | | | |
|-----------------------------|---|---|------------------------------------|---|
| Information | Population and Setting | Intervention/ Study Groups | Definitions | Results |
| Author: Levit ²³ | Number of patients: | Study Groups: | Outcome Definitions: | Primary Outcomes: |
| | N = 2,676 patients | UAC: n=2035 | BSI: CDC/NHSN definition | CLABSI: |
| Year: 2020 | Number of lines: | UVC: n=2017 | | Adjusted rate/ 1000 central-line days: |
| | N= 4,052 lines | Double lumen: n=679 | CLABSI: if no other source was | • aRR: 2.7 (95% CI: 1.1-6.8); P = 0.04 |
| Study | | Single lumen: n=3373 | identified and if the UC was still | Double lumen UVC: 2.0 |
| Design: Cohort | Setting: | | indwelling or had been removed | Single lumen UVC: 0.7 |
| | Level IV NICU | Device/agent: Catheter type; double- | within 48 hours of the onset of | |
| Risk of | | lumen catheter | infection | Cumulative incidence of UVC-related CLABSI |
| Bias: Low | Location: USA | | | • In the first week: <1% |
| | | Standard preventive measures: | Adverse events: | • at day 14: 3.6% |
| | Dates: June 1, | UC insertion is a sterile, bedside | Complications: break/rupture, | • At day 18: 16.5% |
| | 2008 – May 31, 2018 | procedure typically performed by | occlusion, catheter tip | |
| | | advanced practice providers, | malposition, poor perfusion to | BSI: Incidence, n/N (%) |
| | Inclusion Criteria: | pediatric interns and residents, and | lower extremity, CLABSI, | • UAC: 2/2035 (0.1%) |
| | Any infant admitted | neonatal-perinatal medicine fellows | thrombus, or effusion | • UVC: 19/2017 (0.9%) |
| | to the NICU who had | Double-lumen catheter insertion is | | |
| | a UAC, UVC, or both | based solely on anticipated need | Sampling /Testing strategy: NR | Topic-specific outcomes: |
| | successfully placed | Blood is not typically withdrawn | | Mean dwell time, days (range) |
| | (i.e., catheter tip in | from a UVC | Other notes: authors concluded | • UAC: 5.5 days (1-22) |
| | the desired, central | Confirmation of UC placement is via | the risk of CLABSI was low at day | • UVC: 7.6 days (1-21) |
| | location) | thoracoabdominal radiograph | 14 even though the risk increased | |
| | | Routine, scheduled reconfirmation | to 3 times the risk of the first | Adverse events |
| | Exclusion Criteria: | of UC location is not performed | week of life. | All complications: |
| | • NR | Heparin at a concentration of 1 U ml- | | Adjusted rate/ 1000 central-line days |
| | | ¹ of fluid is infused continuously | | • IRR: 0.3 (95% CI: 0.2-0.4) |
| | | through all central line lumens | | • UAC: 4.6 |
| | | Central line tubing utilized for | | • UVC: 17.6 |
| | | parenteral nutrition, intralipids, | | • p = NR |
| | | and/or blood products is changed | | Incidence, n/N (%) |
| | | every 24 hours | | • UAC: 51/2035 (2.5%) |
| | | Tubing utilized only for dextrose | | • UVC: 269/2017 (13.3%) |
| | | containing fluids is changed every 96 | | • p = NR |
| | | hours | | Adjusted rate/ 1000 central-line days |
| | | An assessment of the continued | | • Double lumen UVC: 17.2 |
| | | need for central access is typically | | • Single lumen UVC: 17.2 |
| | | made at day 5-7 of use | | • p = 0.23 |
| | | | | φ - 0.23 |
| | | | | Complications excluding catheter malposition: |
| | | | | Adjusted rate/ 1000 central-line days |
| | | | | • aIRR: 2.3 (95% CI: 1.2-4.6); p = 0.02 |
| | 1 | | | - ann. 2.3 (33/0 Ci. 1.2 7.0), p - 0.02 |

| Study | | | | |
|--|---|---|--|--|
| Information | Population and Setting | Intervention/ Study Groups | Definitions | Results |
| Author: Sanderson ² Year: 2017 Study Design: Multicenter retrospective cohort Risk of Bias: Low | Number of patients: N= 3985 Number of lines: N= 6000 • UVC: 2668 • PICC: 3332 Total catheter days: 43, 302 • Baseline characteristics were significantly different between groups: including Gestational age, birth weight, congenital anomaly, PPROM, respiratory distress, cesarean | Study groups: UVC only: n=1392 UVC only: n=1317 UVC and PICC: n=1276 Standard preventive measures: NR | Outcome Definitions: First CLABSI: CDC 2016 definition and consistent with and within 48 hours of CVC removal (consistent with NSW criteria). CLABSI assigned to CVC in situ. Repeated organism isolates w/in 14 days of LOS diagnosis is not considered new LOS. Early onset sepsis (EOS): positive blood culture in an infant taken within the first 48 h of life and a clinical picture consistent with sepsis. Late onset sepsis (LOS): a positive blood culture, clinical symptoms, and signs of sepsis | Double lumen UVC: 3.8 Single lumen UVC: 1.6 Adjusted rate/ 1000 central-line days IRR: 1.6 (95% CI: 1.02-2.5) UAC: 3.9 UVC: 2.4 p = NR Primary Outcome: CLABSI: Incidence: n (%) UVC: 116/2668 (4.3%) PICC: 287/ 3332 (8.6%) p < 0.01 Rate: n/ 1000 catheter days UVC: 9.88 PICC: 9.09 UVC CLABSI rate: increased beyond 4days, and by days 6-7 had more than 5 times the risk (IRR: 5.85 (1.18-28.96) of CLABSI than on days 45. Topic-specific Outcomes: Dwell time: "The hazard ratio (HR) of UVC and PICC diverged |
| | distress, cesarean delivery, major surgery, mortality, perinatal asphyxia/ trauma, age at first insertion, duration of CVC Setting: Multicenter: 10 NICUs in 10 hospitals Location: Australia Dates: January 1, 2007 – December 31, 2009 Inclusion Criteria: All | | symptoms, and signs of sepsis and clinician decision to treat with antibiotics for 5 days (including CoNS) Sampling /Testing strategy: Blood/catheter tip culture. Other notes: None | beyond the 3-4 days dwell time. UVC had a higher HR and earlier rise than PICC." • "the incremental CLABSI rate increase was highest in UVCs of infants with UVC+PICC, which almost doubled every 2-3 days between days 2 and 7 (14, 27, and 45 per 1000 line-days respectively) and continued to rise with increasing duration, peaking at 85 per 1000 line-days at days 10 and 11." • "the hazard function for CLABSI showed that the group with early PICC insertion (before day 4) had a trend of lower HR." Adverse events: Mortality w/in 14 days of CLABSI (%LOS deaths) • UVC: 8/1392 (61.3%) • PICC: 1/1317 (16.0%) • UVC+PICC: 11276 (5.0%) |
| | infants born during the study dates admitted to 1 | | | • 0VC+PICC: 11276 (5.0%) • p < 0.001 |

| Study | | | | |
|----------------------------|----------------------------------|---|----------------------------------|---|
| Information | Population and Setting | Intervention/ Study Groups | Definitions | Results |
| | of 10 NICUs with one or | , | | |
| | more UVCs or PICCs | | | |
| | inserted. | | | |
| | Exclusion Criteria: NR | | | |
| Author: | Number of patients: | Study groups: | Outcome Definitions: | Primary Outcomes: |
| Vachharajani ³⁴ | N = 201 | Post-QI1: Jan 1, 20014 – March 30, | CLABSI & UVC-associated CLABSI: | CLABSI: |
| | Number of lines: | 2014: introduction of QI initiative | not defined | • Pre-QI: 1 (in situ 8 days) |
| Year: 2017 | N = 201 | including questionnaire, staff education, | | • QI: 2 (in situ for 7 & 10 days) |
| | | and standardization of feeding protocol: | Sampling /Testing strategy: | UVC-associated CLABSI QI to Pre-QI: |
| Study Design: | Setting: NICU, University | Feeding GL for preterm infants: | NR. | • IRR 1.13 (95% CI 0.469 – 2.332); p = 0.92 |
| Uncontrolled | Hospital | BW≤1000g | | 2.25 (55% 6. 6. 1.65 2.552), p |
| before-after | · | Starting volume: 10ml/kg | Other notes: None | Topic-specific outcomes: NR |
| | Location: USA | Advance volume: 10ml/kg during | | UVC> 7days |
| Risk of Bias: | | morning rounds | | • PRE-QI: 23/86 (27%) |
| Moderate | Dates: Jan 1, 2012 – June | When to fortify: 60-100ml/kg | | • QI1: 42/115 (36.5%) |
| | 30, 2014 | BW≥1000g | | • p = 0.045 |
| | | Starting volume: 20ml/kg | | - P - 0.0 13 |
| | Inclusion Criteria: | Advance volume: 20ml/kg during | | Adverse events: NR |
| | uncomplicated NICU | morning rounds | | Autorise events. |
| | patients without | When to fortify: 80-100ml/kg | | |
| | congenital anomalies | Questionnaire implemented to | | |
| | with GA>27 wks. or | encourage providers to consider leaving | | |
| | >1000g at birth, | the existing UVC in situ if neonate met | | |
| | extubated by 3 days of | criteria. Encourage provider to remove | | |
| | age and on enteral feeds | UVC and insert PICC after day 7 if | | |
| | by 2 – 3 days of age | neonate not tolerating 60-70ml/kg/ day | | |
| | | of feeds by 5-6 days of age. | | |
| | Exclusion Criteria: babies | Post QI2: April 1, 2014 – June 30, 2014 | | |
| | who died within a week | Pre-QI: Jan 1, 2012 – December 31, | | |
| | following redirection of | 2013 | | |
| | care. Neonates with | baseline | | |
| | abdominal wall defects, | | | |
| | congenital heart defect, | Standard preventive measures: NR | | |
| | congenital diaphragmatic | | | |
| | hernia, spontaneous | | | |
| | intestinal perforation, | | | |
| | neonates requiring >7d | | | |
| | antibiotic therapy. | | | |
| Author: Butler- | Number of patients: | Patient Groups: | Outcome Definitions: | Primary Outcomes: |
| O'Hara ³³ | N = 986 | Pre-intervention Jan – Oct 2006 | CLABSI: infant was considered | CLABSI: |
| | Number of lines: | Post-intervention: After November | to have a CLABSI when one of | Multiple logistic regression model: |
| Year: 2012 | N = 986 | 2006 | these two criteria were met: (1) | • Year (2006, 2007 vs 2008, 2009) 4.10 (1.29-13.0); p |
| | | | the infant had a recognized | = 0.02 |

| | Population and Setting | Intervention / Study Groups | Definitions | Results |
|--|---|---|---|---|
| Study Design: Uncontrolled before after study (Retrospective cohort) Risk of Bias: Moderate | Population and Setting Setting: Neonatal ICU Location: USA Dates: January 1, 2006 – December 31, 2009 Inclusion Criteria: All infants for whom UVC was placed as part of routine care. Exclusion Criteria: NR | Intervention/ Study Groups Infants >7 days UVC group: n=448 Infants in this group were smaller and had lower gestational age at birth. Infants ≤ 7 days UVC group: n=536 Assess impact of evidence based catheter insertion and maintenance bundle. Multi intervention: November 2006 All providers in NICU in contact with central catheters received education, evidence-based checklists for UVC and PICC insertions, dressing changes, and care and maintenance of UVC and PICC during solution changes. PICC Team: dedicated 4 hours/day exclusively to catheter care and maintenance and changing of central catheter solutions. Team not responsible for umbilical venous or arterial catheter care or fluid changes. Provided care for most but not all days each month. Parenteral nutrition solutions for PICCs were changed once daily. Team used procedure carts specifically for PICC care and maintenance. used a closed medication administration system and adhered to strict evidence-based practices for solution changes and catheter care. hand hygiene and maintained aseptic technique when changing all intravenous tubing and when entering the catheter, including scrubbing the catheter hub with povidone-iodine. | pathogen cultured from one or more culture sites and the organism cultured from the blood was not related to an infection at another site; and (2) the infant had symptoms (eg, fever, hypotension) and positive laboratory results not related to an infection at another site and a common skin contaminant (eg, coagulasenegative staphylococcus) was cultured from two or more blood cultures drawn on separate occasions. Sampling /Testing strategy: Blood and catheter tip cultures performed. Other notes: None | Results ● Birthweight, kg 0.20 (0.02-1.71); p = 0.14 ● Gestational age, weeks 0.92 (0.70-1.20); p = 0.52 ● UVC in place >7 days 5.48 (1.18-25.50); p = 0.03 ● Initial antibiotics >3 days 0.28 (0.10-0.76); p = 0.01 CLABSI Rate/ 1000 days & HR (95% CI) and duration of CVC ≤7 days ● UVC: 1.0; 1 ● PICC: 6.1: 1 8-10 days: ● UVC: 5.4; 5 (0.98 – 51.00) ● PICC: 1.4; 0.2 (0.02 – 1.60) 11-14 days: ● UVC: 21; 20 (5 – 185) ● PICC: 3.8; 0.6 (0.2 – 3.1) >14 days: ● UVC: 9.2; 1.5 (0.6 – 5.8) Topic-specific outcomes: None Adverse events: NR |

| Study | | | | |
|-------------|------------------------|---|-------------|---------|
| Information | Population and Setting | Intervention/ Study Groups | Definitions | Results |
| | | of infection and dressing integrity. PICC | | |
| | | care done by assistant buddy system. | | |
| | | | | |
| | | Standard preventive measures: | | |
| | | UVC Placement care: | | |
| | | care of the umbilical site included use | | |
| | | of betadine for cord preparation before | | |
| | | catheter placement. | | |
| | | No triple dye applied to any umbilical | | |
| | | cord that required a UVC. Either a | | |
| | | single- or double lumen catheter was | | |
| | | inserted in sterile conditions. A second | | |
| | | assistant or "buddy" was assigned and | | |
| | | dedicated to placement of the UVC. | | |
| | | Care of the catheters was standardized, | | |
| | | with use of evidence-based bundled | | |
| | | care and a series of procedural | | |
| | | checklists. Catheters were sutured in | | |
| | | place in the umbilical cord, and tape was then used to secure the catheter to | | |
| | | the infant's abdomen. The clinical team | | |
| | | (not the PICC team) was responsible for | | |
| | | changing the fluids of the umbilical | | |
| | | arterial | | |
| | | and venous catheters. At the | | |
| | | completion of the procedure, a | | |
| | | procedural checklist was completed | | |
| | | indicating use of sterile technique from | | |
| | | the start of the procedure until the final | | |
| | | placement and suture of the catheter. | | |
| | | · | | |
| | | PICC insertion/care: | | |
| | | Placement of the PICC was performed in | | |
| | | sterile conditions. Povidone-iodine | | |
| | | solution swabbed 360 degrees | | |
| | | surrounding the chosen insertion site. | | |
| | | Either a 25- or 30-cm catheter with a | | |
| | | 24-gauge introducer needle was | | |
| | | inserted in the infant's brachial, axillary, | | |
| | | saphenous, or external jugular vein. | | |
| | | Dressings were assessed hourly and | | |
| | | changed when loss of adhesiveness, | | |
| | | drainage at the site, or the dressing | | |

| Study | | | | |
|--|---|---|--|--|
| Information | Population and Setting | Intervention/ Study Groups | Definitions | Results |
| | | became too restrictive. A "second assistant" or "buddy" was available for PICC insertion, dressing changes and maintenance. Dedicated team of performed all dressing changes and catheter manipulations. Checklists were used for PICC insertion, catheter dressing changes, and care and maintenance of the PICC during solution | | |
| • 11 - 5 - 11 | | changes. | 0 | |
| Author: Butler O'Hara ³⁵ | Number of patients: N=210 Number of lines: | Patient Groups: Long term (n=104) UVC was replaced when the catheter was no longer | Outcome Definitions: Catheter related infection: defined infection while a catheter | Primary Outcomes: Catheter related infection rate/ 1000 catheter days: • Long term: 11.5 |
| Year: 2006 | N = 210 | needed or by 28 days at the latest. UVC replaced with PCVCs | (UVC or PCVC) was in place. Each infant was counted only once as | • Short term: 7.4 |
| Study Design: RCT | Setting: Neonatal ICU Location: Boston, | Short term: (n=106) The umbilical venous catheter remained in place up | having a catheter infection during the study regardless of future blood-culture results. | Catheter-related infection Incidence: • Long term: 21/104 |
| Risk of Bias: Low | Massachusetts, USA | to 7 to 10 days of age. If central access was necessary | Sampling /Testing strategy: | Short term: 14/106 OR: 1.66 (95% CI: 0.79 – 3.48); p = 0.17 p = 0.18 |
| | Dates: July 1998 - February 2004 | beyond day 10, PCVC placement was attempted beginning at day 7 to assure | All infants who had a sepsis workup performed during the | Topic-specific outcomes: |
| | Inclusion Criteria: Infants with birth weights ≤1250 g who had a UVC placed on NICU admission. Infants born at <24 weeks' gestation or <500 | successful placement by day 10. Standard preventive measures: ■ Both infusion and flush solutions contained heparin (1.0 IU/ml for infants >1000 g and 0.5 IU/ml for infants ≤1000g or on total | study period (until 28 days or until catheter removal, whichever came first) had simultaneous quantitative peripheral and catheter blood cultures performed. | Catheter duration before infection, days, median: • Long term: 14.0 • Short term: 11.5 • p = 0.35 Adverse events (n) |
| | g at birth, but attending neonatologist was first consulted and had to provide approval. | parenteral nutrition. • Catheters sutured in place into the umbilical cord, and tape was then used to secure the catheter to the | Other notes: None | Thromboses: • Long term: 7 • Short term: 4 Pericardial effusions |
| | Exclusion Criteria: Infants who required a UVC for | infant's abdomen. Placement of PCVC performed under sterile conditions, and care of | | Long term: 10 Short term: 11 |
| | exchange transfusion, infants with gastrointestinal | catheters was standardized. • The catheter and the proximal | | NEC (Bell's 40 stage 2 or above) • Long term: 11 |
| | abnormalities including gastroschisis and | portion of the extension set were secured to the skin by using a sterile, transparent, occlusive | | Short term:7 Mortality: Long torm: 7 |
| | omphalocele, or infants with congenital heart | dressing. | | • Long term: 7 • Short term: 8 |

| Study | | | | |
|------------------------|-----------------------------|---|---------------------------------|---|
| Information | Population and Setting | Intervention/ Study Groups | Definitions | Results |
| | disease with intracardiac | Solution infusing through the PCVC | | Arrhythmia |
| | shunting. | contained heparin (at the same | | Long term: 1 |
| | | concentrations as for UVC) and ran | | • Short term: 0 |
| | | at a minimum rate of 1.0 ml/hour. | | Embolus |
| | | Sterile gloves were worn during all | | None observed |
| | | solution changes. | | Hemorrhage |
| | | Intravenous tubing was secured well | | None observed |
| | | to the skin but did not occlude any | | Pleural effusion |
| | | part of the dressing. | | None observed |
| | | Dressing integrity was assessed | | Liver disease (one-year follow-up) |
| | | routinely and documented. | | • Long term: 1 |
| | | Dressings were changed when there | | Short term: 0 |
| | | was loss of adhesiveness or drainage | | Broken catheter |
| | | at the site or when they became too | | None observed |
| | | restrictive. | | Catheters removed due to mechanical complications |
| | | | | • Long term: 27/181 |
| | | | | • Short term: 27/210 |
| Author: | Number of patients: | Patient groups: | Outcome Definitions: | Primary Outcomes: |
| Bhandari ¹² | N = 2091 | Patients: n = 2091 | Nosocomial sepsis: Presence of | Total Nosocomial Sepsis: % infected was significantly |
| | Number of lines: | | clinical signs of infection, | different for each catheter type: P<0.0001 |
| Year: 1997 | N = 2091 | Standard preventive measures: | initiation of anti-microbial | Umbilical artery |
| | | UA and UV were placed either by | therapy and a positive blood | • Infected: 179/1699 (10.5%) |
| Study Design: | Setting: 2 NICUs, 1 at a | the physicians or the neonatal nurse | culture obtained from a | • Non-infected: 1520/ (89.5%) |
| Prospective | University Hospital, 1 at a | practitioners (NNP) at both the | peripheral site or via the | Umbilical venous: |
| cohort study | regional hospital | NICUs. | catheter after the third | • Infected: 81/617 (13.1%) |
| | | • Tunneled CVs (Broviac) were placed | postnatal day. | • Non-infected: 536/617 (86.9%) |
| Risk of Bias: | Location: USA | by pediatric surgeons | | Central Venous |
| High | | Percutaneous central venous | Association between duration of | • Infected: 99/294 (33.5%) |
| | Dates: Regional Hospital | placements were done exclusively | catheter use, type, and | • Non-infected: 194/294 (66.2%) |
| | November 11, 1987 - | by the NNPs using a standard | nosocomial sepsis at University | Percutaneous Catheter |
| | December 31, 1993 | protocol (sterile technique and site | hospital: the incidence of | • Infected: 96/308 (31.2%) |
| | | preparation with povidone iodine) | positive blood cultures from | • Non-infected: 212/308 (68.8%) |
| | University Hospital: | Some PCVs placed as "long | time of insertion of catheter | Peripheral Artery |
| | January 1, 1989 - | peripheral" lines rather than as | until 3 days after removal was | • Infected: 35/189 (18.5%) |
| | December 31, 1993 | central lines for technical reasons. | analyzed for a consecutive | Non-infected: 154/189 (71.5%) |
| | | Catheter maintenance was done per | population subset over 2.5 | ,, |
| | Inclusion Criteria: All | nursing protocols at both hospitals: | years | Nosocomial Sepsis and Dwell Time: n (%) |
| | neonates admitted to the | sterile dressing and IV tubing | | Umbilical artery |
| | 2 hospital NICUs if one or | changes. | Infants with bacteremia: | • 1-3 days: 1/207 (0.5%) |
| | more vascular catheter | Peripheral arterial catheters were | - And >1 catheter | • 4-7 days: 4/175 (2.3%) |
| | was simultaneously or | placed by physicians/NNPs | simultaneously: each | • 8-14 days: 7/62 (11.3%) |
| | sequentially placed: | All lines had heparin infusions. | | • ≥15 days: 4/19 (21.1%) |

| Study | | | | |
|-------------|----------------------------|----------------------------|---|-----------------------------|
| Information | Population and Setting | Intervention/ Study Groups | Definitions | Results |
| | umbilical artery (UA), | | catheter was included in | • ≥8 days: 13.6% |
| | Umbilical venous (UV), | | analysis for association | • ≤7 days: 1.3% |
| | central venous Broviac | | - And >1 catheter sequentially: | • p < 0.0001 |
| | (CV), percutaneously | | the last catheter place was | Umbilical venous: |
| | placed central venous | | assigned the infection. | • 1-3 days: 1/129 (0.8%) |
| | (PC), or peripheral artery | | - 1/3 of infants with CV or PC | • 4-7 days: 4/58 (6.9%) |
| | (PA). | | compared 10-18% of infants | • 8-14 days: 3/52 (5.8%) |
| | | | with other catheter types. | • ≥15 days: 1/5 (20.0%) |
| | Exclusion Criteria: NR | | Committee /Testine stretery | Central Venous |
| | | | Sampling /Testing strategy: Blood/catheter tip culture. | • 1-3 days: 0/4 (0%) |
| | | | Biood/catheter tip culture. | • 4-7 days: 1/6 (16.7%) |
| | | | Other notes: Incidence of | • 8-14 days: 2/30 (6.7%) |
| | | | infection by comparing different | • ≥15 days: 14/72 (19.4%) |
| | | | catheter types. | Percutaneous Catheter |
| | | | dutileter types. | • 1-3 days: 0/12 (0%) |
| | | | | • 4-7 days: 0/13 (0%) |
| | | | | • 8-14 days: 1/27 (3.7%) |
| | | | | • ≥15 days: 3/27 (11.1%) |
| | | | | Peripheral Artery |
| | | | | • 1-3 days: 1/30 (3.3%) |
| | | | | • 4-7 days: 0/27 (0%) |
| | | | | • 8-14 days: 1/9 (11.1%) |
| | | | | • ≥15 days: 0/3 (0%) |
| | | | | |
| | | | | Topic-specific outcomes: NR |
| | | | | Adverse events: NR |

Table 45 Risk of Bias for Randomized Controlled Trials on Umbilical Catheter Dwell Times

| Author Year | Described as randomized | Randomization appropriately performed | Described as double-blind | Outcome assessor blinded | Study participant blinded | Investigator blinded | Attrition described | Attrition smaller than 10-15% of assigned patients | Attrition appropriately analyzed | Funding source(s) disclosed and no obvious conflict of interest | Overall Risk of Bias |
|---|-------------------------------|---------------------------------------|---------------------------|--------------------------------|---------------------------------|-------------------------|---------------------|---|--|---|-------------------------|
| Butler O' Hara 2006 ³⁵ | ✓ | ✓ | | | √ | | √ | ~ | ~ | ~ | Low |

. Page **81** of **137**

Table 46 Risk of Bias for Two Group Studies on Umbilical Catheter Dwell Times

| Author Year | Were patients randomly assigned to the study's groups? | For non-randomized trials, did the study employ any other methods to enhance group comparability such as matching, stratification, or statistical methods to adjust for baseline differences? | Did patients in different study groups have similar levels of performance on the outcome of interest and other important factors at the time they were assigned to groups? | Did the study enroll all suitable patients or consecutive suitable patients within a time period? | Was the comparison of interest prospectively planned? | Were the two groups treated/ evaluated concurrently? | Was the study blinded or double- blinded? | • | Risk of Bias |
|--------------------------|--|---|--|---|---|--|---|------------|-----------------|
| | groups: | baseline unterences? | groups: | a time perious | piailileur | concurrently | billided: | directions | DIdS |
| Levit 2020 ²³ | ✓ | ✓ | ✓ | ✓ | ✓ | ✓ | ✓ | | Low |

Table 47 Risk of Bias for Single Group Studies on Umbilical Catheter Dwell Times

| Author Year | Did the study enroll all suitable patients or consecutive suitable patients within a time period? | • | Was the funding for this study derived from a source that would not benefit financially from results in a particular direction? | Risk of Bias |
|------------------------------------|---|---|---|-----------------|
| Bhandari 1997 ¹² | ✓ | ✓ | | High |
| Sanderson 2017 ² | ✓ | ✓ | ✓ | Moderate |
| Vachharajani 2017 ³⁴ | ✓ | ✓ | ✓ | Moderate |

Table 48 Risk of Bias for Two Group Studies on Umbilical Catheter Dwell Times

| Author Year | Were patients randomly assigned to the study's groups? | For non-randomized trials, did the study employ any other methods to enhance group comparability such as matching, stratification, or statistical methods to adjust for baseline differences? | Did patients in different study groups have similar levels of performance on the outcome of interest and other important factors at the time they were assigned to groups? | Did the study enroll all suitable patients or consecutive suitable patients within a time period? | Was the comparison of interest prospectively planned? | Were the two groups treated/ evaluated concurrently? | Was the study blinded or double-blinded? | Was the funding for this study derived from a source that would not benefit financially from results in a particular direction? | Risk of Bias |
|---|--|---|--|---|---|--|--|---|-----------------|
| Butler- O'Hara 2012 ³³ | | ✓ | ✓ | ✓ | ✓ | ✓ | | | Moderate |

C.10. Optimal Peripherally Inserted Central Catheter Dwell Time

Key Question 10. What is the optimal duration for peripherally inserted central catheters to prevent CLABSI in NICU patients?

Table 49 Summary of Findings on Peripherally Inserted Central Catheter Dwell Times to Prevent CLABSI

| , | or manage of the representation of the first to the first | Quantity and Type of | GRADE of Evidence for Outcome |
|---------------------------|--|--|---|
| Outcome | Findings | Evidence | and Limitations of the Evidence |
| CLABSI* | Three observational studies^{2, 36, 37} reported increasing risk of CLABSI with increasing PICC dwell time, but no clear inflection point for PICC removal or replacement to reduce CLABSI risk. One observational study² found that increasing dwell time was associated with increased risk of CLABSI for PICCs, but reported no clear inflection point for PICC removal or replacement. One observational study³⁶ reported the risk of CLABSIs increased during the 2 weeks after PICC insertion and then remained elevated until PICC removal but data did not point to a clear inflection point beyond which infection increases. One observational study³⁷ reported an increase in CLABSI risk of 14% per day between catheter days 1-18, and of 33% per day from days 35 through 60. One observational study⁷ reported that compared with the risk of CLABSI in week 1, no other week was associated with increased risk of CLABSI for PICCs suggesting no clear optimal PICC dwell time to reduce CLABSI risk. | 4 OBS n=3332 PICCS ² n=4797 PICCS ³⁶ n=683 PICCS ³⁷ n=14,451 PICCS ⁷ | Low |
| Catheter-related BSI* | One observational study³⁸ reported increasing dwell time was a significant factor for the odds of developing CRBSI (p<0.01), however the optimal timing for removal of a PICC could not be determined. One observational study³⁹ reported that for each week of PICC duration, the trend was for an increasing rate over time; however, this did not reach significance (p = 0.09) and dwell time was not a predictor of the odds of developing CR-BSI. (OR: 1.19 (0.91–1.57); p = 0.212). Almost all PICCs in this study were removed within 2 weeks after insertion. One observational study⁴⁰ found no difference in the mean dwell time between infected and non-infected patients. (p = 0.6064). | 3 OBS N=412 PICCS ³⁸ N=946 PICCS ³⁹ N=63 PICCS ⁴⁰ | Low |
| Catheter –related sepsis* | One observational study⁴¹ found the odds of developing CRS was 3 times higher if the catheter was in place for ≥9 days (OR: 3.1 (95% CI: 1.64-5.87); p<0.01). | 1 OBS n=294 PICCS ⁴¹ | Very Low • Imprecision: only one study |

Table 50 Extracted Information on Peripherally Central Catheter Dwell Time

| Study | Population and | Intervention/ Study | | |
|------------------------|------------------|-----------------------|---|--------------------------|
| Information | Setting | Groups | Definitions | Results |
| Author: | Number of | Patient group: | Outcome Definitions: | Primary Outcomes: |
| Sanderson ² | patients: | UVC only: n=1,392 | First CLABSI: CDC 2016 definition and consistent with | CLABSI: |
| | N = 3,985 | UVC only: n=1,317 | and within 48 hours of CVC removal (consistent | Incidence: n (%) |
| Year: 2017 | Number of lines: | UVC and PICC: n=1,276 | with NSW Health criteria*). CLABSI assigned to CVC | • UVC: 116/2668 (4.3%) |
| | n=6,000 | | in situ. Repeated organism isolates w/in 14 days of | • PICC: 287/ 3332 (8.6%) |
| | • UVC: 2,668 | | LOS diagnosis is not considered new LOS. | |

| Study | Population and | Intervention/ Study | | |
|---|--|----------------------------------|--|--|
| Information | Setting | Groups | Definitions | Results |
| Study Design: Multicenter retrospective | PICC: 3,332Total catheterdays: 43, 302Baseline | Standard preventive measures: NR | * available at: http://www.cec.health.nsw.gov.au/ data/assets/pdf_file/0009/258372/hai- | p < 0.01Rate: n/ 1,000 catheter daysUVC: 9.88 |
| cohort Risk of Bias: | characteristics were significantly | | manual.pdf Early onset sepsis (EOS): positive blood culture in an infant taken within the first 48 hrs. of life and a | PICC: 9.09 UVC CLABSI rate: increased beyond 4 days, and by days 6-7 group 1 [UVC only] had more than five |
| Low | different among groups (UVC only [group 1], PICC only [group 2], UVC and PICC [group 3]): including gestational age, birthweight, congenital anomaly, PPROM, respiratory distress, | | clinical picture consistent with sepsis. Late onset sepsis (LOS): a positive blood culture, clinical symptoms, and signs of sepsis and clinician decision to treat with antibiotics for ≥5 days (including CoNS) Incidence of CLABSI: expressed as number of episodes per 1,000 catheter-days and number of episodes per 1,000 catheters inserted PPROM: prolonged premature rupture of membranes IRR: incidence rate ratio Sampling /Testing strategy: Blood/catheter tip culture. | times the risk (IRR: 5.85 (CI: 1.18-28.96) of CLABSI than on days 45. Dwell time: "The hazard ratio (HR) of UVC and PICC diverged beyond the 3-4 days dwell time. UVC had a higher HR and earlier rise than PICC." "the incremental CLABSI rate increase was highest in UVCs of infants with UVC+PICC, which almost doubled every 2-3 days between days 2 and 7 (14, 27, and 45 per 1,000 line-days respectively) and continued to rise with increasing duration, peaking at 85 per 1,000 line-days at days 10 and 11." "the hazard function for CLABSI showed that the group with early PICC insertion (before day 4) had a trend of lower HR." |
| | cesarean delivery, major surgery, mortality, perinatal asphyxia/ trauma, age at first insertion, duration of CVC Setting: Multicenter: 10 NICUs in 10 hospitals Location: Australia | | Other notes: None | Topic-specific outcomes: NR Adverse events: Mortality w/in 14 days of CLABSI (% LOS deaths) • UVC: 8/1,392 (61.3%) • PICC: 1/1,317 (16.0%) • UVC+PICC: 1/1,276 (5.0%) • p < 0.001 |

Page 84 of 137

| Study | Population and | Intervention/ Study | | |
|------------------------|--|---|--|---|
| Information | Setting | Groups | Definitions | Results |
| | Dates: January 1, | - | | |
| | 2007 – December | | | |
| | 31, 2009 | | | |
| | | | | |
| | Inclusion Criteria: | | | |
| | All infants born | | | |
| | during the study | | | |
| | dates admitted to | | | |
| | 1 of 10 NICUs with | | | |
| | one or more UVCs | | | |
| | or PICCs inserted. | | | |
| | | | | |
| | Exclusion Criteria: | | | |
| | NR | | | |
| Author: | Number of | Patient group: | Outcome Definitions: | Primary Outcomes: |
| Greenberg ⁷ | infants: | N = 13,327 NICU infants | CLABSI: NHSN 2008 definition. | CLABSI: |
| | N = 13,327 | | Positive blood culture for a recognized pathogen | Incidence |
| Year: 2015 | Number of lines: | Tunneled catheters | not related to an infection at another site | Tunneled catheters: 39/1,116 (3.5%) |
| | N = 15,567 | (n= 1,116/15,567; 7.2 %)) | Diagnosis of CLABSI required systemic signs and | • PICCs: 199/ 14,451 (1.4%) |
| Study | Catheter days: | | symptoms of infection and isolation of the same | • p <0.001 |
| Design: | N = 256,088 | PICCs | organism from ≥ 2 blood cultures drawn on | Rate |
| retrospective | | (n = 14,451/15,567; 93%) | separate occasions. | • 0.93 CLABSI / 1,000 catheter days |
| cohort study | Setting: | | CLABSI attribution: | |
| _ | Multicenter NICU | Device/agent: Catheter | If a single catheter had multiple associated positive | Effect of dwell time on CLABSI |
| Risk of Bias: | (141 NICUs; 13 | type | blood cultures (occurred on 12 occasions), only the | Week 1 |
| Low | states) | | first positive blood culture was included in the | • Tunneled catheters: 5/1,116 (0.4%) |
| | | Standard preventive | analysis. | HR (95% CI:) reference |
| | Location: USA | measures: | If a CLABSI occurred in the presence of multiple | • PICCs: 82/14,451 (0.6%) |
| | Data a Cantanahan | Participating sites adopted | catheters (this occurred on 3 occasions), the CLABSI | HR (95% CI): reference |
| | Dates: September | a central catheter insertion | was attributed to both catheters. | Week 2 |
| | 2011 – August | and maintenance bundle | Dwell time: number of days from line insertion until | • Tunneled: 5/969 (0.5%) HR: 1.3 (0.4 – 4.4) |
| | 2013 | which included: | either line removal or day of CLABSI. The day of line | • PICCs: 56/8,250 (0.7%) |
| | Inclusion Criteria: | Hygiene for insertion | insertion was defined as line day 1; weeks of dwell | • HR 1.2 (95% CI: 0.9 – 1.7) |
| | | Daily assessment of line | time were categorized into 7-day periods starting on | Week 3 |
| | Infant with PICCs or | need | line day 3 (week 1 = line days 3–9, week 2 = line days | • Tunneled: 3/748 (0.4%) HR: 1.0 (0.2 – 4.4) |
| | tunneled | A recommendation to | 10–16, etc.). | • PICCs: 31/4,061 (0.8%); HR 1.3 (0.8 – 1.9) |
| | catheters | remove central lines when infants achieved | A diverse avente. ND | Week 4 |
| | obtained from | | Adverse events: NR | • Tunneled: 2/580 (0.3%) HR: 0.9 (0.2 – 4.7) |
| | NCLABSI | 120 mL/kg per day of | Someting /Testing stretomy Bland sultimes | • PICCs: 5/2,209 (0.2%); HR 0.4 (0.1 – 0.9) |
| | database | enteral feedings | Sampling /Testing strategy: Blood cultures | Week 5 |
| | during study | techniques for sterile | Other meters | • Tunneled: 3/452 (0.7%) HR: 1.8 (0.4 – 7.6) |
| | dates | dressing changes and | Other notes: HR: hazard ratio | • PICCs: 7/1,290 (0.5%); HR 0.9 (0.4– 1.9) |
| | dutes | catheter access. | TIN. Hazdiu Idliu | Page 85 of 137 |

| Study | Population and | Intervention/ Study | | |
|----------------------|------------------------------------|--------------------------------------|--|---|
| Information | Setting | Groups | Definitions | Results |
| | | Antibiotic practices | | Week 6 |
| | Exclusion Criteria: | were not standardized | | • Tunneled: 4/355 (1.1%) HR: 3.2 (0.8 – 12.0) |
| | Central lines | between the sites. | | • PICCs: 7/765 (0.9%); HR 1.5 (0.7– 3.2) |
| | inserted and | | | Week 7 |
| | removed | | | • Tunneled: 4/280 (1.4%); HR 4.0 (1.1-15.4) |
| | within the first | | | • PICCs: 4/453 (0.9%); HR 1.4 (0.5-4.0) |
| | 2 days | | | Week 8 |
| | Positive blood | | | • Tunneled: 1/288 (0.4%); HR 1.3 (0.1-11.4) |
| | cultures | | | • PICCs: 3/278 (1.1%); HR 1.6 (0.5-5.2) |
| | occurring | | | Week 9 |
| | within 2 days | | | • Tunneled: 3/178 (1.7%); HR: 4.7 (1.1-20.3) |
| | of line | | | • PICCs: 2/183 (1.1%); HR: 1.5 (0.4-6.3) |
| | placement | | | Week 10 |
| | | | | • Tunneled: 1/151 (0.7%); HR: 2.0 (0.2-17.7) |
| | | | | • PICCs: 0/125 (0) |
| | | | | V 1 1003. 0/ 123 (0) |
| | | | | Topic-specific outcomes: |
| | | | | Catheter dwell time median, (IQR) |
| | | | | • Tunneled catheters: 24.5 d (14-45) |
| | | | | • PICCs: 11 d (7-18) |
| | | | | • p < 0.001 |
| | | | | β < 0.001 |
| | | | | Adverse events: NR |
| Author: | Number of | Patient group: N = 63 | Outcome Definitions: | Primary Outcomes: |
| Rangel ⁴⁰ | patients: | | Catheter-related Infection: categorized as positive or | Catheter-related infection: |
| · · | N = 63 | Standard preventive | negative according to the result of the blood culture | Positive Blood Culture: 16/63 (25.40%) |
| Year: 2014 | Number of lines: | measures: | | |
| | N = 63 | A protocol for the | Sampling /Testing strategy: | Topic-specific outcomes: |
| Study | | insertion and | Blood culture. | Indwell Time mean (SD), days |
| Design: | Setting: NICU, 1 | maintenance of PICC | | Catheter-related infection: 10.69 (± 6.322) |
| Retrospective | university hospital | lines, | Other notes: None | • No infection: 9.88 (± 4.87) |
| cohort study | | A routine for recording | | • p = 0.6064 |
| | Location: Brazil | procedures undertaken | | |
| Risk of Bias: | | with the PICC by the | | Adverse events: NR |
| Moderate | Dates: January | nursing professionals in | | |
| | 2009 - December | a surveillance form for | | |
| | 2010 | intravascular devices | | |
| | | filed in the medical | | |
| | Inclusion Criteria: | records, | | |
| | NICU newborns | A technical body | | |
| | weighing | trained and | | |

Page **86** of **137**

| Study | Population and | Intervention/ Study | | |
|-----------------------------------|----------------------------------|---------------------------|--|--|
| Information | Setting | Groups | Definitions | Results |
| ormacion | 500 - 1,499 g, | empowered for the use | Deminions . | incounts |
| | born in the | of this type of protocol. | | |
| | institution | of this type of protocol. | | |
| | between | | | |
| | | | | |
| | January 2009 - December 2010, | | | |
| | with a record of | | | |
| | having had a PICC | | | |
| | | | | |
| | line in that period. | | | |
| | Exclusion Criteria: | | | |
| | NICU newborns | | | |
| | with congenital | | | |
| | malformations, | | | |
| | diagnosis of | | | |
| | infection prior to | | | |
| | the implantation | | | |
| | of the PICC, who | | | |
| | were suspected of | | | |
| | primary | | | |
| | bloodstream | | | |
| | infection (BSI) or | | | |
| | who were | | | |
| | transferred due to | | | |
| | any situation were | | | |
| | excluded from the | | | |
| | study. | | 2.0 | |
| Author: Milstone ³⁶ | Number of | Patient group: N= 3,967 | Outcome Definitions: | Primary Outcomes: |
| wiistones | patients: N = 3,967 | Standard preventive | PICC dwell time: days from PICC insertion until either PICC removal or the date of CLABSI, whichever was | Catheter-related sepsis: PICC-associated CLABSI, incidence, n/N (%): 149/4,797 |
| Year: 2013 | Number of lines: | measures: | earlier. | (3.1%) |
| Teal. 2013 | N = 4,797 PICCs | Trained infection | PICC-associated CLABSI: CDC 2008 NHSN definition of | PICC-associated CLABSI incidence rate/1,000 days: 1.66 |
| Study | Number of | preventionists | CLABSI occurring in a PICC | Time from PICC insertion to CLABSI, median (range), |
| Design: | catheter days: | performed prospective | "two or more blood cultures drawn on separate | days: 18 (1–166) |
| Retrospective | N = 89,946 | surveillance to monitor | occasions" for common skin commensal bacteria (i.e., | days. 10 (1 100) |
| cohort | 11 - 03,340 | positive blood cultures | coagulase negative staphylococci | CLABSI Incidence rate/ 1,000 catheter days (95% CI) |
| 3011011 | Setting: | in patients with | - SSSSSSSSSSSSSSSSSSSSSSSSSSSSSSSSSSSS | • 1-10d: 1.05 (95% CI: 0.77–1.41) |
| Risk of Bias: | multicenter; NICU | indwelling catheters by | Sampling /Testing strategy: | • 11-20d: 1.98 (95% CI: 1.44–2.66) |
| Moderate | (8), university | using laboratory | Blood/catheter tip culture. | • 21-30d: 2.07 (95% CI: 1.31–3.11) |
| | hospitals | databases and infection | , | • 31-40d: 2.47 (95% CI: 1.38–4.07) |
| | | surveillance support | Other notes: | • 41-50d: 1.73 (95% Cl: 0.63–3.76) |
| | Location: USA | systems | IRR: incidence rate ratio | • 51-60d: 2.95 (95% CI: 1.08–6.41) |
| | | | | - 31 000. 2.33 (33/0 Cl. 1.00-0.41) |
| | | | | Dogs 97 of 127 |

| Study | Population and | Intervention/ Study | | |
|--|--|---|---|--|
| Information | Setting | Groups | Definitions | Results |
| Information | Dates: January 1, 2005- June 30, 2010 Inclusion Criteria: Neonates who had a PICC inserted in a NICU during the study dates. Exclusion Criteria: NR | Groups | Median PICC dwell time of 14 days; 25% remained in place for ≥ 23 days | **Nesuts • >60d: 3.31 (95% CI: 1.65–5.92) • "PICCs w/ dwell time of 8 - 13 days, 14 – 22 d, and ≥23 days each had an increased risk of infection compared w/ PICCs in place for ≤7 days" (p <0.05). • "there is no clear inflection point after which the daily risk of CLABSIs increases" **Topic-specific outcomes:* PICC dwell times, n (%) • ≤7 d:1,096 (22.9) • 8–13 d: 1,289 (26.8) • 14–22 d: 1,129 (23.6) • ≥23 d 1,283 (26.7) Univariate analysis: Catheter dwell time: CLABSI (%), unadjusted IRR (95% CI); p • ≤7 d: 25 (16.6%), 1.0 (reference) • 8–13 d: 32 (21.2%), 2.02 (1.21–3.38); p = 0.007 • 14–22 d: 39(25.8%), 3.27 (2.04–5.24); p < 0.001 • ≥23 d: 55(36.4%), 2.71 (1.71–4.27); p < 0.001 |
| Author: Ohki ³⁹ Year: 2013 Study Design: Prospective cohort study | Population: N = 946 Number of lines: N = 946 Setting: Multicenter NICU (19) Location: Japan | Patient group: N=946 Number of lines: n=946 PICCs Standard preventive measures: Institution insertion practices were classified into three groups: 1) Those with MRP (i.e. | Outcome Definitions: CR-BSI: one of the following signs or symptoms: fever (>38°C), hypothermia (<36°C), apnea, or bradycardia, plus at least one positive blood culture from a patient with a PICC, without an infection at another site. PICC- associated BSI: if the line was in use during the preceding 48 hr. period. Extremely low-birthweight (ELBW): birthweight <1000 g Very low-birthweight (VLBW): birthweight <1500 g, | Adverse events: NR Primary Outcomes: Catheter-related BSI: Duration of PICC (per each 1 week) Multivariate analysis: OR: 1.19 (95% CI: 0.91–1.57) p = 0.212 Topic-specific outcomes: NR Adverse events: NR |
| Risk of Bias: Moderate | Dates: February 2005 - March 2007. Inclusion Criteria: Neonates >21 weeks of gestational age, | 1) Those with MBP (i.e., cap, mask, sterile gown, sterile gloves, and large sterile drapes: MBP group), 2) Those with standard barrier precautions (i.e., sterile gloves and small | PCE/CT: determined by ultrasonography. Pleural effusion/ascites: identified on ultrasonography or standard radiography. Catheter removal difficulties: inability to remove the catheter after local warming or local massage, and requirement for procedures such as guidewire reinsertion or surgical removal. | |

| Study | Population and | Intervention/ Study | | |
|---------------------|---------------------|--|--|--|
| Information | Setting | Groups | Definitions | Results |
| | weighing >400 g | sterile drape: SBP group), | Symptomatic catheter-related thrombosis: thrombosis | 11000111 |
| | at birth, and | and | seen on venography or ultrasonography and | |
| | without lethal | 3) Those that conducted | associated with clinical symptoms. | |
| | congenital | the procedure similarly | Asymptomatic catheter-related thrombosis: excluded | |
| | anomalies or | to peripheral line | from analysis because routine ultrasonography was | |
| | major | placement (i.e., without | conducted at only two institutes. | |
| | chromosomal | preparing a sterile field, | , | |
| | abnormalities. | the operator pulls the | Sampling /Testing strategy: Blood culture. | |
| | | catheter from the vinyl | | |
| | Exclusion Criteria: | sheath with small sterile | Other notes: None | |
| | Patients | forceps, and inserts it | | |
| | transported from | from the introducer | | |
| | study institutions | needle without touching | | |
| | with a PICC in situ | the PICC: non-PICC | | |
| | | group) | | |
| Author: | Number of | Patient group: N=218 | Outcome Definitions: | Primary Outcomes: |
| Njere ⁴¹ | Patients: | Number of lines: n=294 | Catheter-related sepsis: positive blood cultures | Catheter-related sepsis: |
| | N = 218 | PICC lines | (peripheral/central) and/or a positive tip culture | Rate/ 1,000 catheter days: 17 (21%) |
| Year: | Number of lines: | | after removal in the presence of a clinical suspicion | Odds of infection: |
| 2011 | N = 294 | Standard preventive | of line sepsis. | Catheter in situ ≥9 days: OR: 3.1 (95% CI: 1.64-5.87); |
| | | measures: | Sepsis: in the presence of a catheter, the patient | p<0.01 |
| Study | Setting: Neonatal | Insertion: | developed temperature instability, tachypnea, | Multivariable analysis included dwell time, incubator |
| Design: | ICU; tertiary | Aseptic technique: use of | apnea, lethargy, and abdominal distension, a rising | vs. open crib, catheter type, previous infected line, |
| Prospective | referral hospital | sterile set, theater | C-reactive protein, or nonspecific factors. | number of previous lines, attempts at insertion & |
| cohort | Location: UK | gowns, gloves, drapes, | PICC line infection: positive peripheral or central blood culture or a positive catheter tip culture after removal | gestational age. |
| Risk of Bias: | Location. OK | catheters, and other equipment. Use of masks | in the presence of clinical signs of catheter-related | Only significant predictor: of PICC line infection: |
| Moderate | Dates: January | and caps was not | sepsis | dwell time ≥9 days |
| Wioderate | 2006 to June 2009 | considered an essential | 364313 | Topic-specific outcomes: |
| | 2000 to Julic 2003 | part of aseptic technique. | Sampling /Testing strategy: | CONS isolated from blood culture: 55/62 (89%). |
| | Inclusion Criteria: | Skin prep: chlorhexidine | Blood/catheter tip culture. | CONSTSORATED HOTH BIOOD CUITURE. 33/02 (83/0). |
| | Neonates who | gluconate 0.05% and | | Adverse events: |
| | had PICCS for | allowed to dry. | Other notes: None | Reasons for catheter removal |
| | parenteral | , | CONS: coagulase-negative staphylococcus | Possible infection: 77/ (20.2%) |
| | nutrition and | Catheter care: | | Leakage/extravasation: 45/294 (15.3%) |
| | venous access. | Run saline when not in | | Blocked: 4/ (1.4%) |
| | | use (not heparinized) | | |
| | Exclusion Criteria: | Catheters accessed | | |
| | Incomplete data | after washing hands, | | |
| | on Neonate | donning sterile gloves, | | |
| | | cleaning connector | | |
| | | hubs with .05% CHG, | | |
| | | and allowing to dry. | | |

| Study | Population and | Intervention/ Study | | |
|-------------------|---|---|--|--|
| Information | Setting | Groups | Definitions | Results |
| | | Secured with Steristrips | | |
| | | and occlusive | | |
| | | transparent dressings | | |
| | | Dressing replacement: | | |
| | | removed if loose and | | |
| | | new dressing reapplied. | | |
| | | Tubing Change: every | | |
| | | 24hrs when parenteral | | |
| _ | | nutrition bags changed | • | |
| Author: | Number of | Patient group: N=275 | Outcome Definitions: | Primary Outcomes: |
| Hsu ³⁸ | patients: | VLBW infants | CRBSI: At least one positive blood culture obtained | CRBSI: |
| | N = 275 | PICCs: n=412 | from a peripheral vein, the presence of clinical | • Episodes: 67/412 (16.3%) |
| Year: 2010 | Number of lines: | PICC lines | features consistent with bloodstream infection in | Rate/ 1000 catheter days: 8.3 |
| | N = 275 | | the presence of a PICC in position, and no other site | Time from placement to CRBSI: 16.4 ± 8.4 days |
| Study | | Standard preventive | of infection. | Multivariable logistic regression including Dwell time, |
| Design: | Setting: Neonatal | measures: | Phlebitis: when a linear red streak developed along the | insertion site, birthweight, gestational age, weight. |
| Retrospective | ICU | Insertion: | superficial veins from the insertion site. | Duration of PICC: p<0.01 (Area under curve 0.68) |
| cohort study | Lasatiana Tairran | Under sterile | Thrombosis: suspected when leg swelling with or | • Femoral insertion site: OR: 1.76, 95% CI: 1.01-3.07; p |
| Risk of Bias: | Location: Taiwan | environment by nursing | without poor perfusion developed. | < 0.045 |
| | Datas Isasani | specialist or | Catheter site inflammation: diagnosed in the presence | |
| Moderate | Dates: January 2005 to December | residents/fellows under | of lymphangitis, purulence, or at least two signs of inflammation (erythema, tenderness, increased | Univariate analysis: |
| | 2005 to December 2006 | supervision | warmth, or induration). | Duration of PICC, days; case no/total no, incidence (%) |
| | 2000 | Vein selected by those | Cholestasis: direct bilirubin ≥ 1.5 mg/dL. | • ≤10 days: 6/92; 6.2%) (reference) |
| | Inclusion Criteria: | who performed catheter insertion and | Rupture: completely broken PICCs, rather than simple | • 11-20 days: 10/98, 10.2%); RR: 1.72, 95% CI: 0.60- |
| | Very low | peripheral veins | leakage. | 4.94 |
| | birthweight | preferred over femoral | Extravasation: dislodgement of PICC. | • ≥21: days: 51/217 (23.5%) RR: 4.66, 95% CI: 1.93- |
| | (VLBW) infants | vein. | Time to complication: calculated from day of insertion | 11.28 |
| | admitted to the | Skin disinfection: | to day recognition of any catheter-related | Site of insertion, incidence (%) |
| | NICU with a | rubbing the site of | complication. | • Non-femoral: 30/241 (12.4%) |
| | percutaneously | insertion with sterile | | • Femoral: 37/171 (21.6%) |
| | inserted catheter | gauze soaked in a | Sampling /Testing strategy: | |
| | inserted into a | solution of 10% PI | Blood culture. | Topic-specific outcomes: NR |
| | central vein | containing 75% alcohol. | | (b) (b) (c) 1 (d) |
| | | The same disinfectant | Other notes: | Adverse events: incidence, n/N (%); rate/1000 catheter |
| | Exclusion Criteria: | applied to insertion site | No bacterial pathogens were identified from blood | days |
| | percutaneous | after successful | cultures for both phlebitis and catheter site | • Phlebitis: 25/412 (6.1%); 3.1/1,000 catheter days |
| | catheters inserted | insertion; saline used to | inflammation. | • Thrombosis: 1/412 (0.2%); 0.12/1,000 catheter days |
| | into non-central | decolorize and covered | | • Catheter site inflammation: 28/412 (6.8%); 3.5/1000 |
| | veins | by transparent | | catheter days |
| | | dressing. | | • Leakage: 7/412 (1.7%); 0.9/1,000 catheter days |
| | | Maintenance: | | • Rupture: 10/412 (2.4%); 1.2/1,000 catheter days |
| | | | | • Extravasation: 4/412 (1.0%); 0.5/1,000 catheter days |

| Study | Population and | Intervention/ Study | | |
|---------------------------|---------------------------------------|--|--|--|
| Information | Setting | Groups | Definitions | Results |
| | 3 | Manipulations | | • Occlusion: 32/412 (7.8%); 4.0/1,000 catheter days |
| | | performed using | | |
| | | standard protocol by | | |
| | | NICU nurses. | | |
| | | Decision for PICC | | |
| | | removal made by | | |
| | | neonatologist or senior | | |
| | | resident; phlebitis, | | |
| | | catheter fracture, | | |
| | | extravasation, | | |
| | | thrombosis and | | |
| | | catheter site | | |
| | | inflammation were | | |
| | | definitive indications | | |
| | | for removal and | | |
| | | infected catheters | | |
| | | always removed with | | |
| | | positive cultures or | | |
| | | infant unresponsive to | | |
| | | IV antibiotics | | |
| Author: | Population: N= | Patient group: | Outcome Definitions: | Primary Outcomes: |
| Sengupta ³⁷ | 683 | N = 683 NICU patients with | CLABSI: CDC/NHSN 2002 Guideline definition | CLABSI: |
| | | PICC | | Incidence/ PICC n/N (%): 21/683 (3.1%) CLABSI |
| Year: 2010 | PICC lines = 953 | | PICC: peripherally inserted central venous catheter that | Incidence (over study period): |
| | | PICC lines: 917/953 eligible | terminates at or close to the heart or in 1 of the great | 2.01/1,000 catheter days; (95% CI: 1.24-3.06) PICC |
| Study | Setting: NICU at | for analysis | vessels and is used for infusion, withdrawal of blood, or | associated CLABSI |
| Design: | tertiary care | | hemodynamic monitoring | |
| Retrospective | hospital | Standard preventive | DIGG CARGE CARGE CARGE | Topic-specific Outcomes: |
| cohort study | | measures: | PICC associated CLABSI: primary bloodstream infection | PICC duration: |
| Diels of Dies: | Location: | PICCs placed by designated | in a patient admitted to the NICU for > 48 hrs. before | (interval, no. of events, incidence) |
| Risk of Bias: Moderate | US | trained nurse or physicians Standard protocol followed | the onset of infection that met the NHSN criteria for CLABSI | 1-10 days = 6; 1.08/1,000 catheter days |
| iviouerate | Dates: Jan 1, | re insertion and | PICC follow-up time(duration): days from line insertion | 11-20 days = 8; 2.77/1,000 catheter days |
| | 2006-Dec 31, | maintenance practices | until 1 of the following: | 21-30 days = 4; 2.7/1,000 catheter days |
| | 2008-Dec 31, | As part of a quality | 1) date of CLABSI, | 31-40 days = 0 |
| | 2000 | improvement initiative to | 2) termination of the PICC, or | 41-50 days = 1; 2.29/1,000 catheter days |
| | Inclusion Criteria: | reduce CLABSI, hospital | 3) administrative censoring at discharge from the | 51-60 days = 2; 7.78/1,000 catheter days |
| | Eligible patients | epidemiology and infection | NICU | Univariate analysis of PICC as risk factor for CLABSI: |
| | had a PICC | control dept. monitors | Only the first CLABSI was included for a patient who | (days since PICC insertion, IRR, 95% CI) |
| | inserted in the | development of | had multiple CLABSIs from the same PICC | < 19 days: IRR = 1.15 (1.05-1.26) |
| | NICU between Jan | bacteremia in patients | | p < 0.01 |
| | 1, 2006-Dec 31, | | Sampling /Testing strategy: Blood culture | 19-35 days: IRR = 0.80 (0.67-0.96) |
| | 2008. In patients | | , 0, 0 | p = 0.02 |
| | a a a a a a a a a a a a a a a a a a a | I | | Dags 01 of 127 |

| Study | Population and | Intervention/ Study | | |
|-------------|---------------------|---------------------|-------------------|---|
| Information | Setting | Groups | Definitions | Results |
| | with multiple | | Other notes: None | > 35 days: IRR = 1.32 (1.12-1.55) |
| | PICCs, only the | | | p = < 0.01 |
| | first was included | | | Multivariable analysis of PICC as risk factor for CLABSI: |
| | in analysis | | | (days since PICC insertion, IRR, 95% CI) |
| | | | | < 19 days: IRR = 1.14 (1.04-1.25) |
| | Exclusion criteria: | | | p = < 0.01 |
| | PICCs terminated | | | 19-35 days: IRR = 0.80 (0.66-0.96) |
| | the same day | | | p = 0.02 |
| | inserted and PICCs | | | > 35 days: IRR = 1.33 (1.12-1.57) |
| | removed within | | | p = < 0.01 |
| | 48 hrs. of NICU | | | |
| | admission | | | Adverse Events: NR |
| | excluded | | | |

Page **92** of **137**

Table 51 Risk of Bias for Two Group Studies on Percutaneous Central Catheter Dwell Times

| Author Year | Were patients randomly assigned to the study's groups? | For non-randomized trials, did the study employ any other methods to enhance group comparability such as matching, stratification, or statistical methods to adjust for baseline differences? | Did patients in different study groups have similar levels of performance on the outcome of interest and other important factors at the time they were assigned to groups? | Did the study enroll all suitable patients or consecutive suitable patients within a time period? | Was the comparison of interest prospectively planned? | Were the two groups treated/ evaluated concurrently? | Was the study blinded or double- blinded? | Was the funding for this study derived from a source that would not benefit financially from results in a particular direction? | Risk of Bias |
|--------------------------------|--|---|--|---|---|--|---|---|-----------------|
| Greenburg 2015 ⁷ | ✓ | | ✓ | ✓ | ✓ | ✓ | ✓ | ✓ | Low |
| Sanderson 2017 ² | ✓ | | ✓ | ✓ | √ | ✓ | | ✓ | Low |

Table 52 Risk of Bias for Single Group Studies on Percutaneous Central Catheter Dwell Times

| Author Year | Did the study enroll all suitable patients or consecutive suitable patients within a time period? | Was the study prospectively planned? | Were independent or blinded assessors used to assess subjective Outcome Definitions, or were the Outcome Definitions objective? | Was the funding for this study derived from a source that would not benefit financially from results in a particular direction? | Risk of Bias |
|--------------------------------|---|--------------------------------------|---|---|--------------|
| Hsu 2010 ³⁸ | ✓ | | ✓ | | Moderate |
| Milstone 2013 ³⁶ | ✓ | | ✓ | | Moderate |
| Njere 2011 ⁴¹ | ✓ | | ✓ | | Moderate |
| Ohki 2013 ³⁹ | ✓ | ✓ | ✓ | | Moderate |
| Rangel 2014 ⁴⁰ | ✓ | ✓ | ✓ | | Moderate |
| Sengupta 2010 ³⁷ | ✓ | | ✓ | | Moderate |

C.11. Dedicated Catheter Care Team

Key Question 11. In NICU patients requiring central catheters, does the use of dedicated catheter care teams compared with standard of care, prevent CLABSI?

Table 53 Summary of Findings for a Dedicated Percutaneous Inserted Central Catheter Care Team vs. Standard_of Care to Prevent CLABSI

| | | Quantity and Type | |
|---------|--|---------------------------|-------------------------------|
| | | of Evidence | GRADE of Evidence for Outcome |
| Outcome | Findings | (Sample Size) | (Limitations of the Evidence) |
| | • 1 single center OBS study ⁴² implemented a central line maintenance team in the NICU and | 1 OBS | Very Low |
| CLABSI* | reported a significant decrease in overall CLABSI rates comparing pre- and post-line team | n=NR lines ⁴² | Imprecision: only one study |
| | rates [11.6 vs. 4.0 per 1000 catheter days, P<0.001]. | | |
| | 1 single center OBS study⁴³ implementing dedicated vascular access team in NICU | 1 OBS ⁴⁴ | Very Low |
| | reported no difference in CRBSI rates for all indwelling lines [23/100 (23%) vs. 24/100 | n=200 lines ⁴³ | Imprecision: only one study |
| CRBSI* | (24%); $p = 0.868$]; however, a duration stratification analysis revealed a 49% reduction in | | |
| CRESI | CRBSI for indwelling PICC lines ≥30 days: 39/47 (83%), p = 0.0407; no difference for | | |
| | indwelling lines $<$ 30 days: short (0-3 days): 2/47 (4.3%), p = NS; intermediate (4-29 days): | | |
| | 6/47 (12.8%), p = NS. | | |

Table 54 Extracted Information on a Dedicated Percutaneous Inserted Central Catheter Care Team

| Study Information | Population and Setting | Intervention/ Study Group | Definitions | Results |
|------------------------|--------------------------------|------------------------------------|---|------------------------------------|
| Author: Holzmann- | Number of patients: | Intervention: | Outcome Definitions | Primary Outcomes: |
| Pazgal ⁴² | N = NR | Catheter care team: | CLABSI | CLABSI, rate/ 1000 line day (after |
| | Number of lines: | Recruitment: Sixteen bedside | CDC-2004 National Healthcare Safety | correcting for NHSN definition |
| Year: 2012 | N = NR | nurses and seventeen neonatal | Network (NHSN) definitions. Definition | change and excluding skin |
| | | transport nurses | changed 2008 | contaminants): |
| Study Design: Before- | Setting: Level III to III NICU | Education & Training: intensive | | • Pre-intervention: 11.6 |
| after study | | education repeated on evidence- | | • Intervention: 4.0 |
| | Location: US | based practices for central line | Sampling /Testing strategy: NR | • p < 0.001 |
| Risk of Bias: Moderate | | management already in place in | | · |
| | Dates: December 2006 – | the unit. Training utilized | Other notes: None | Weight-specific CLABSI, rate/ |
| | September 2010 | standardized written protocols | | 1000 line days: |
| | | developed by infection control and | | <750g |
| | Inclusion Criteria: NR | NICU nursing leadership that | | Pre-intervention: 15.6 |
| | | formalized established guidelines | | • Intervention: 6.1 |
| | Exclusion Criteria: NR | for performance maintenance | | • p = 0.012 |
| | | Line maintenance: tubing changes, | | |
| | | dressing changes, and accessing of | | 751-1000g |
| | | central lines for blood draws or | | Pre-intervention: 9.7 |
| | | medication administration. Every | | • Intervention: 5.3 |
| | | member of the line team had to | | • p = 0.095 |
| | | learn proper procedures and | | , |
| | | techniques for line maintenance, | | 1001-1500g |

| | | perform the procedure while being observed by a trainer and be checked off upon satisfactory | | Pre-intervention: 12.8 Intervention: 3.2 |
|------------------------------|--|--|---|--|
| | | demonstration of competence. March 2008, the line team took over performance of all tubing changes, accessing of central lines for blood draws and all dressing changes. Line team members worked in teams of two to perform dressing changes and tubing changes. Only members of the line team could perform these functions on any central line. October 2009: line team took over medication administration through central lines, however in Control: Pre-Intervention: December 2006 – | | p = 0.001 1501-2500g Pre-intervention: 9.8 Intervention: 2.1 p = 0.001 >2500g Pre-intervention: 9.5 Intervention: 2.5 p < 0.001 Topic-specific outcomes: NR Adverse events: NR |
| | | March 2008, baseline Device/agent: Central care team Monitoring intervention: NA Standard preventive measures: NR | | |
| Author: Taylor ⁴³ | Number of patients: | Intervention: | Outcome Definitions | Primary Outcomes: |
| 2,12 | N = 200 | PICC team: n = 100 | Catheter-related bloodstream infection | CRBSI, n/N (%): |
| Year: 2011 | Number of lines: | | (CRBSI): | • Pre-intervention: 23/100 |
| | N = 200 | April 14, 2006 | Positive blood culture with recognized | (23%) |
| Study Design: | | Percutaneously inserted central | pathogen, or | • Intervention: 24/100 (24%) |
| Prospective cohort | Setting: Level IIIC NICU | catheters (PICC) team established that included neonatal nurse | positive blood culture with common skin contaminant or positive antigen test on | • p = 0.868 |
| Risk of Bias: Low | Location: US Dates: Pre-intervention: March 1, 2005-March 31, 2006; Post-intervention (PICC team): June 22, 2006-July 9, 2007 Inclusion Criteria: All extremely low birth weight infants (≤1000g) admitted to a level IIIC | practitioners, neonatology fellows, NICU transport nurses, and selected NICU bedside nurses. Policies established for early patient identification for line placement, regular surveillance of line site and dressing integrity, and tracking of complications Standardized training developed according to national guidelines to | blood and temperature instability (>100.4°C), hypotension, apnea or bradycardia, and • Signs and symptoms with positive laboratory results not related to infection at another site (e.g., necrotizing enterocolitis) Short duration: central lines between 0-3 days | Survival analysis (attributable to CRBSI): • Hazard ratio: 0.48 (95% CI: 0.25-0.91) • p = 0.025 CRBSI, patients with short central line duration (0-3 days), n/N (%): • 2/47 (4.3%) |

Exclusion Criteria: Infants born in the 2-week period when the PICC team was being formulated.

improve aseptic precautions, promote best practice, and to minimize variability in technique among team members.

A formalized system developed for tracking weekly, and as necessary dressing changes for all and lines, including chlorhexidine patches

PICC dressing changes and line assessments performed weekly; daily line changes are the responsibility of the bedside registered nurse.

Control:

Pre-intervention: n=100

Incoming neonatology fellows, transport nurses, and neonatal nurse practitioners would receive bedside training for PICC placement by their senior peers.

Dressing changes would be performed by fellows, transport nurses, and nurse practitioners on an as needed basis, with the goal of once per week.

Patients needing PICC lines identified when bedside nurse would approach the medical team for intravenous access or when it was noted that an umbilical line needed to be replaced (14-day maximum).

Documentation of PICC placement or removal was done via a free-text procedure note in the medical record. No set system for documentation or tracking of dressing changes, although date of last dressing change was kept in a log

Intermediate duration: central lines between 4-29 days

Sampling /Testing strategy: Blood cultures performed.

Other notes: It is acknowledged that some infants in the control group were exposed toward the end of their hospitalization to the benefits of the PICC team if they were still hospitalized after the PICC team was established. However, given the direction of these differences, it is most likely that any such effect would have led to an underestimation of the intervention-related reduction in CRBSI risk.

April 2005

Adopted the closed medication system

CRBSI, patients with intermediate central line duration (4-29 days), n/N (%):

• 6/47 (12.8%)

CRBSI, patients with highest central line duration (≥30 days), n/N (%):

- 39/47(83%)
- 49% reduction
- p = 0.0407

Topic-specific outcomes:

Time to CRBSI, median (range):

- Pre-intervention: 30 (5-70)
- Intervention: 35 (1-82)
- p = 0.360

Central line days, median (range):

- Pre-intervention: 7 (0-100)
- Intervention: 18 (1-141)
- p = 0.009

Adverse events:

Mortality (not attributable to CRBSI), n/N (%):
Pre-intervention: 15/100 (15%)

Intervention: 27/100 (27%)

p = 0.056

maintained by the on-service neonatology fellow. March 2006 Didactic and clinical training to improve aseptic precautions, promote "best practice," and minimize variability to technique among team members were completed (continued an ongoing basis for new members). After a 2-hr. didactic training session, new team members demonstrated proficiency by completing PICC insertions and dress changes under the guidance of a preceptor. Device/agent: NA Monitoring intervention: NA Standard preventive measures: Sterile prep for PICC placement was done with full sterile gown, mask, gloves and 10% iodine solution. Dressing changes were done with mask and sterile gloves, using 2% chlorhexidine swabs. Dressing changes included replacement of chlorhexidine dressing for infants older than 30 days or 32 weeks.

Table 55 Risk of Bias for Two Group Studies on a Dedicated Percutaneous Inserted Central Catheter Care Team

| Author Year | All study groups derived from similar source/reference populations | Attrition not significantly different across study groups | Measure of exposure is valid | Measure of outcome is valid | Investigator blinded or were outcomes well-defined and objective to endpoint assessment | Potential confounders identified | Statistical adjustment for potential confounders done | Funding source(s) disclosed and no obvious conflict of interest | Overall Risk of Bias |
|---|--|---|------------------------------|-----------------------------|---|----------------------------------|---|--|----------------------------|
| Holzmann- Pazcal 2012 ⁴² | ✓ | | ✓ | ✓ | | ✓ | | | Moderate |
| Taylor 2011 ⁴³ | √ | ✓ | √ | √ | √ | √ | ✓ | | Low |

C.12. Central Line Insertion and Maintenance Bundles

Question 12. In NICU patients that are the optimal elements of central line insertion and maintenance bundles to prevent CLABSI?

Table 56 Summary of Findings on Insertion and Maintenance Bundles vs. Standard of Care to Prevent CLABSI

| Outcome | Findings | Quantity and Type of Evidence (Sample Size) | GRADE of Evidence for Outcome (Limitations of the Evidence) |
|--|---|---|--|
| CLABSI* | • Three observational studies ⁴⁵⁻⁴⁷ reported a reduction of CLABSI rate. | 3 OBS N=NR ⁴⁵ N=NR ⁴⁶ N=NR ⁴⁷ | Low |
| Healthcare Personnel Bundle Compliance* | • Three observational studies ⁴⁵⁻⁴⁷ reported increases in compliance with bundle elements. | 1 OBS ⁴⁵ N=NR N=NR ⁴⁶ N=NR ⁴⁷ | Low |

Table 57 Extracted Information for Central Venous Catheter Insertion and Maintenance Bundles

| Study | Population and | Intervention/ Study | | |
|-----------------------------|----------------------|------------------------|--|--------------------------------|
| Information | Setting | Groups | Definitions | Results |
| Author: Balla ⁴⁷ | Number of patients: | Patient Groups: n=229 | Outcome Definitions: | Primary Outcomes |
| | N = 229 | Number of lines: n=229 | BSI: A laboratory-confirmed bloodstream infection that | CLABSI rate per 1000-line days |
| Year: 2018 | Number of lines: N = | Baseline: n = 54 | was not secondary to an infection at another site. | Baseline: 31.74 |
| | 229 | • 3 months | CLABSI: A primary BSI in a patient that had a central line | • Phase 1: 18.58 |
| Study Design: | Catting NICI | Intervention: n = 175 | within the 48-hour period | • Phase 2: 3.73 |
| Interrupted time series | Setting: NICU | • 12 months | before the development of the BSI was considered CLABSI. | • Phase 3: 3.53 |

| Study | Population and | Intervention/ Study | | |
|---------------------------|---|--|--|---|
| Information | Setting | Groups | Definitions | Results |
| Information Risk of Bias: | Location: USA Dates: June 2015 – August 2016 Inclusion Criteria: All patients (aged 0 months to 21 years) admitted to the hospital who received a central line, as defined by the NHSN, comprised the study population. The NHSN defines a central line as an intravascular catheter that terminates at or close to the heart or 1 of the great vessels and is used for infusion, withdrawal of blood, or hemodynamic monitoring. Exclusion Criteria: Exclusion of a patient from the study occurred only if the patient had received a central line before admission and developed a bloodstream infection within 48 hours of admission with supporting clinical or laboratory evidence of an infection at the time of admission. This exclusion criterion is | Surveillance Denominator data collection: A monthly roster for denominator data collection displayed on the QI board was successful. Audits of the denominator data were performed on 5 random days per month to verify the accuracy. Hand hygiene: Change in HH policy: revised from routine hand wash to hand rub. Education & training: All the HCPs were educated about HH through posters, regular classes and one to one communication. Performance & Feedback Sharing data regularly during monthly ward meetings, giving feedback both group and individualized, including personnel from all levels of care in the team Compliance assessment: The compliance with HH was studied with the help of audits, which found that the main problem was duration of hand | Compliance Indicators: The process indicators were based on hand hygiene (30 audits per month) and central line care audits (10 audits per month). • If all the steps of hand hygiene including the six core steps and the duration were correctly performed, it was considered 'overall compliant to HH'. • Central line bundle: The central line care audits focused on insertion practices (number of central lines inserted by eligible Healthcare Personnel (HCP), checklist analysis) and maintenance practices (breaks in circuit, 2 HCP handling the central line, scrubbing the hub for 15 seconds, 2% chlorhexidine used for scrub, use of single lumen central line and needleless connectors). Compliance: Random auditing of at least 10% of lines on each unit by staff nurse CLABSI-prevention champions ensured bundle compliance and evaluated necessity of the line. Sampling /Testing strategy: NR Other notes: None | BSI rate per 1000-line days Baseline: 7.3 Phase 1: 4.6 Phase 2: 4.2 Phase 3: 2.3 Mortality Baseline: 2.9% Intervention: 1.7% Topic-specific outcomes: Compliance with maintenance bundle (%) Baseline: NA Phase 1: 59% Phase 2: 68.2%% Phase 3: 66.7% Adverse events: NR |

| Study | Population and | Intervention/ Study | | |
|-------------|---------------------------------|---------------------------------------|-------------|---------|
| Information | Setting | Groups | Definitions | Results |
| | in line with NHSN | hygiene. The successful | | |
| | definitions issued by | PDSA cycle was to do | | |
| | the Centers for | the hand rub by the | | |
| | Disease Control and | clock for 20-30 seconds. | | |
| | Prevention | It was ensured that a | | |
| | (CDC). | clock with a second | | |
| | Blood cultures that | hand was easily visible | | |
| | were positive on | from each bed of the | | |
| | admission and those reported as | unit. | | |
| | contaminants were | | | |
| | not included. | Designated HCP for | | |
| | not morated. | insertion: | | |
| | | Only those HCPs | | |
| | | certified by the QI team | | |
| | | (those who had assisted | | |
| | | five central line | | |
| | | insertions) were | | |
| | | privileged to place the | | |
| | | central line. A senior | | |
| | | nurse or doctor | | |
| | | supervised the process | | |
| | | of insertion using a | | |
| | | checklist and any | | |
| | | deviation from the | | |
| | | policy was noted and | | |
| | | stopped promptly. | | |
| | | Initially | | |
| | | • Insertion had to be a 2- | | |
| | | person job | | |
| | | Insertion Checklist: | | |
| | | Required but elements | | |
| | | not reported | | |
| | | | | |
| | | Maintenance bundle: | | |
| | | Central line card | | |
| | | displayed on infant | | |
| | | warmer to document | | |
| | | the need of line daily | | |
| | | and number of circuit | | |
| | | breaks; | | |

| Study | Population and | Intervention/ Study | | |
|----------------------|-------------------------------------|---|---|--|
| Information | Setting | Groups | Definitions | Results |
| | | Break in circuit – 2 HCP | | |
| | | iob; | | |
| | | • Scrub the hub – 2% | | |
| | | chlorhexidine for 15 | | |
| | | | | |
| | | seconds | | |
| | | Removal bundle | | |
| | | Review the need every | | |
| | | day and remove as | | |
| | | soon as possible. | | |
| | | Control/Comparison: NA | | |
| | | | | |
| | | Device/agent: NA | | |
| | | Monitoring intervention: | | |
| | | Insertion and maintenance | | |
| | | compliance | | |
| | | Standard preventive | | |
| | | measures: NR | | |
| Author: | Number of patients: | Patient Groups: n=NR | Outcome Definitions: | Primary Outcomes |
| Savage ⁴⁶ | N = NR | Number of lines: n=NR | CLABSI: NR | NICU CLABSI rate per 1000-line days ± SD; p- |
| | Number of lines: N = | Study Periods: | Compliance: Random auditing of at least 10% of lines on | value = compared with preintervention |
| Year: 2018 | NR | Preintervention: 2006 - | each unit by staff nurse CLABSI-prevention | period)): |
| | | 2008 | champions ensured bundle compliance and | Preintervention period: 4.84 ± 1.16 |
| Study Design: | Setting: NICU | • Peri-intervention: 2008 | evaluated necessity of the line. | Peri-intervention period: 2.20 ± 1.11; |
| Interrupted time | | - 2011 | | • p = 0.003 |
| series | Location: USA | Post-intervention: | Sampling /Testing strategy: NR | Post-intervention period: 0.41 ± 1.30 |
| Dial. of Di | Datas: 2000 2044 | February 2011 - | Other material Authorise and Justical a | • p < 0.001 |
| Risk of Bias: | Dates: 2006-2014 | December 2012 | Other notes: Authors conducted a | • 2 nd Peri-intervention period: 0.79 ± 1.27 |
| Moderate | La alcada a Catharda All | • 2 nd Peri-intervention: | root cause investigations utilizing the event-specific | • p < 0.001 |
| | Inclusion Criteria: All | 2013 - 2014 | focus groups as well as a special focus group aimed at | F 13.001 |
| | patients (aged 0 | 2013 2014 | identifying | NICU VLBW CLABSI rate per 1000-line days ± |
| | months to 21 years) admitted to the | | common potential causes. Through this process they identified that the NICU was failing to consistently clean | SD; p-value = compared with preintervention |
| | hospital who | Hospital-wide CLABSI | and disinfect | period)): |
| | received a central | Bundle implemented | patient positioning devices on a daily and as-needed | • Pre-intervention period: 7.55 ± 2.23 |
| | line, as defined by | June 2008 - 2011 | basis. The focus groups also identified that wrist and | Peri-intervention period: 3.41 ± 2.12 |
| | the NHSN, comprised | First peri-intervention | hand jewelry, and hair not kept up and away from the | • p = 0.020 |
| | the study population. | period | face by staff were potential sources of bacteria. Family | • Post-intervention period: 0.72 ± 2.49 |
| | The NHSN | 2008 | and staff noncompliance with hand | • p < 0.001 |
| | THE INTIDIA | | and stail noncompliance with hallu | - h < 0.001 |

| Study | Population and | Intervention/ Study | | | |
|-------------|----------------|---------------------|---|--|--|
| Information | Setting | Groups | Definitions | Results | |
| • | | | hygiene principles, especially after cellular telephone use, and lack of coordination with respiratory therapy and lab blood collection to minimize central line accesses potentially contributed to the increase in CLABSIs. | Pesults • 2 nd Peri-intervention period: 1.00 ± 2.44 • p < 0.001 NICU NLBW CLABSI rate per 1000-line days ± SD; p-value = compared with preintervention period)): • Preintervention period: 1.95 ± 0.96 • Peri-intervention period: 0.84 ± 0.91 • p = 0.232 • Post-intervention period: 0.01 ± 1.07 • p = 0.021 • 2 nd Peri-intervention period: 0.66 ± 1.05 • p = 0.180 CLABSI rate per 1000-line days, (n/N): • Preintervention period: 5.14 (45/8763) • SIR: 1.78; p<0.05 • Peri-intervention period: 2.18 (21/9622) • SIR: 1.30 • Post-intervention period: 0.36 (2/5562) • SIR: 0.29; p<0.05 • 2 nd Peri-intervention period: 0.87 (5/5730) • SIR: 0.78 Topic-specific outcomes: Compliance for entire Hospital • 2013 and 2016: 94% - 99%. Compliance to the maintenance bundle, • 2015: 79% • 2016: 91% Reasons for compliance deviation: • Improper documentation of line necessity • Late dressing changes, or • Administration set tubing changes Adverse events: NR | |

| Study | Population and | Intervention/ Study | | |
|-------------|----------------|---|-------------|---------|
| Information | Setting | Groups | Definitions | Results |
| | | all line interactions and | | |
| | | standardized dressing | | |
| | | change protocol | | |
| | | PICU and medical | | |
| | | floors: 24-h | | |
| | | administration sets and | | |
| | | needleless component | | |
| | | changes for lipids and | | |
| | | blood product and 96 h | | |
| | | for nonlipids | | |
| | | • NICU: 96-h | | |
| | | administration set | | |
| | | tubing change for all | | |
| | | fluids/solutions except | | |
| | | lipids and blood draws. | | |
| | | Lines used for lipids and | | |
| | | blood draws remain at | | |
| | | 24-h change | | |
| | | Administration set | | |
| | | hub/access site cap | | |
| | | change after each | | |
| | | blood draw in all units | | |
| | | except NICU: | | |
| | | Disinfection of patient | | |
| | | area at each shift in | | |
| | | NICU and PICU, | | |
| | | disinfection includes all | | |
| | | items used in the | | |
| | | immediate area of the | | |
| | | patient, such as bed | | |
| | | (including linen), | | |
| | | bedside table, overbed | | |
| | | tables, IV pump, | | |
| | | feeding pumps, diaper | | |
| | | scales, and bedside | | |
| | | supply cabinets | | |
| | | 2011 | | |
| | | Closed system for UAC | | |
| | | in NICU (Figure S1) | | |
| | | | | |

| Study | Population and | Intervention/ Study | | |
|-------------|----------------|---|-------------|---------|
| Information | Setting | Groups | Definitions | Results |
| | | Second peri-intervention | | |
| | | period | | |
| | | 2013 | | |
| | | Monthly rotation and | | |
| | | terminal cleaning of | | |
| | | bedside supply cabinets | | |
| | | in NICU to ensure | | |
| | | Cleanliness of supplies | | |
| | | and cabinets used with | | |
| | | long-term-stay infants. | | |
| | | PICU cleans and | | |
| | | Disinfects cabinet at | | |
| | | least monthly and at | | |
| | | discharge | | |
| | | NICU dressing changed | | |
| | | when loose, wet, or | | |
| | | compromised; all other | | |
| | | units maintain 7-d | | |
| | | dressing change | | |
| | | Umbilical cord cleaned | | |
| | | with CHG before and | | |
| | | after line removal | | |
| | | Exposed PICC lines | | |
| | | removed after another | | |
| | | line established. No | | |
| | | manipulation of line to | | |
| | | insert back under skin | | |
| | | 2014 | | |
| | | • CHG daily body wipe for | | |
| | | children older than age | | |
| | | 2 mo in PICU following | | |
| | | SPS | | |
| | | Recommendations. | | |
| | | Daily linen changes re- | | |
| | | emphasized The unit | | |
| | | time out included | | |
| | | checking patient | | |
| | | identification and | | |
| | | announcing the | | |
| | | procedure, the type of | | |

| Study | Population and | Intervention/ Study | | |
|-------------|----------------|---|-------------|---------|
| Information | Setting | Groups | Definitions | Results |
| | | line to be inserted, and | | |
| | | the site of line insertion | | |
| | | All supplies required | | |
| | | available at bedside | | |
| | | before insertion | | |
| | | Inserter and assistant | | |
| | | use maximal sterile | | |
| | | barrier precautions (i.e., | | |
| | | mask, cap, gown, sterile | | |
| | | gloves, and full body | | |
| | | drape) | | |
| | | Face mask worn by | | |
| | | those within 3 feet of | | |
| | | sterile field | | |
| | | Perform skin antisepsis | | |
| | | with povidone-iodine, | | |
| | | CHG, or alcohol | | |
| | | Skin preparation agent | | |
| | | completely dry at time | | |
| | | of first skin puncture | | |
| | | Procedure stopped if | | |
| | | anyone notes sterility | | |
| | | compromised | | |
| | | Catheter maintenance | | |
| | | checklist: | | |
| | | Volume of infant | | |
| | | feedings in mL/kg per | | |
| | | day | | |
| | | Central lines be | | |
| | | discontinued when | | |
| | | an infant's enteral | | |
| | | feedings reached | | |
| | | 120 mL/kg per day | | |
| | | Daily assessment of | | |
| | | catheter need: | | |
| | | "Do we need the | | |
| | | line today?" | | |
| | | "If there was no line | | |
| | | in place today, | | |
| | | would we place | | |
| | | one?" | | |

| Study | Population and | Intervention/ Study | | |
|-------------|----------------|---|-------------|---------|
| Information | Setting | Groups | Definitions | Results |
| | | Dressing integrity and | | |
| | | site cleanliness | | |
| | | assessed (daily at | | |
| | | minimum) | | |
| | | Dressing and site care if | | |
| | | dressing change | | |
| | | performed | | |
| | | Site cleansed with an | | |
| | | appropriate solution | | |
| | | (povidone-iodine, CHG, | | |
| | | or alcohol) | | |
| | | Cleansing solution | | |
| | | allowed to air-dry | | |
| | | completely | | |
| | | Use of a closed system: | | |
| | | closed system | | |
| | | maintained for infusion, | | |
| | | blood draws, and | | |
| | | medication | | |
| | | administration; closed | | |
| | | system is one in which | | |
| | | entries are made | | |
| | | through needleless | | |
| | | connectors or hubs that | | |
| | | have been disinfected | | |
| | | before use | | |
| | | For all catheter | | |
| | | entries/access | | |
| | | • Scrub needleless | | |
| | | connector or hub | | |
| | | using friction with | | |
| | | alcohol or CHG for | | |
| | | ≥15 seconds | | |
| | | Allow surface of | | |
| | | connector or hub to | | |
| | | dry before entry | | |
| | | Staff wear clean | | |
| | | gloves when | | |
| | | accessing or | | |
| | | entering catheter (if | | |
| | | not using closed | | |
| | | system) | | |
| | | | | |

| Study | Population and | Intervention/ Study | | |
|------------------------------|-----------------------------------|--|---|---|
| Information | Setting | Groups | Definitions | Results |
| | | Control/Comparison: NA | | |
| | | | | |
| | | Device/agent: NA | | |
| | | | | |
| | | Monitoring intervention: | | |
| | | Insertion and maintenance | | |
| | | compliance | | |
| | | Standard preventive | | |
| | | measures: NR | | |
| Author: Fisher ⁴⁵ | Number of patients: | Patient Groups: n=NR | Outcome Definitions: | Primary Outcomes |
| , tatilott i isner | N=NR | Number of lines: n=1308 | CLABSI: used the Centers for Disease Control and | CLABSI rate per 1000-line days, adjusted mean |
| Year: 2013 | Number of lines: | | Prevention, National Healthcare Safety Network | rate: |
| | N=NR | Catheter insertion | definition (June 2008, available at | Pre-intervention: 3.94 |
| Study Design: | | checklist: | https://doi.org/10.1016/j.ajic.2008.03.002) | Post-intervention (through July 2010): 1.16 |
| Prospective | Setting: 13 NICUs | Perform hand hygiene | ,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,, | • Reduction rate: 71% |
| cohort study | | before insertion | Process measures: elements of the insertion and | • p = 0.01 |
| | Location: USA | Unit time out before | maintenance bundles | p 5.62 |
| Risk of Bias: | | procedure | | |
| Moderate | Dates: | The unit time out | Sampling /Testing strategy: NR | CLABSI, n: Intervention: 57 |
| | Pre-intervention | included checking | | |
| | (NHSN data, 10/13 | patient | Other notes: No baseline data for process measures | CLABSI rate per 1000-line days, quarterly |
| | NICUs): January 2008-September | identification and | | (values estimated from fig 3): |
| | 2008-September 2009 | announcing the | Compliance measures were limited to 9 points. Statistical | January 2008: 4.6 |
| | 2009 | procedure, the type of line to be | process control (SPC) guidelines suggest a minimum of | April 2008: 5.2 |
| | Intervention (NHSN | inserted, and the | 12 data points to determine significant changes in | July 2008: 3.1 October 2008: 4.0 |
| | data, 13/13 NICUs): | site of line All | control limits on the basis of trends of \$7 points, but that | October 2008: 4.0 |
| | October 2009-June | supplies required | would not limit our ability to detect signals of change and draw conclusions. | January 2009: 3.3 |
| | 2010 | available insertion | and draw conclusions. | April 2009: 5.1 |
| | | At bedside before | Baseline data from 10/13 reported sites; 3/13 level II | July 2009: 3.8 |
| | Post-intervention: | insertion | sites reported no infections based on NHSN criteria from | October 2009: 2.2 |
| | One quarter after | Inserter and assistant | January 2008 through September 2009 | |
| | intervention, and one | use maximal sterile | | January 2010: 2.0 |
| | year later, July- | barrier precautions (i.e., | | April 2010: 1.1 |
| | September 2011 | mask, cap, gown, sterile | | July 2010: 0.9 |
| | Inclusion Criteria: | gloves, and full body | | |
| | Perinatal Quality | drape) | | July 2011: 0.5 |
| | Collaborative of | Face mask worn by | | 13/13 NICHa shawada wa duattar ta CLARCI |
| | North Carolina | those within 3 feet of | | 12/13 NICUs showed a reduction in CLABSI |
| | (PQCNC) invited all | sterile field | | rates |
| | hospitals in the state | | | Topic-specific outcomes: |

| Study | Population and | Intervention/ Study | | |
|-------------|------------------------|---|-------------|---|
| Information | Setting | Groups | Definitions | Results |
| | with a NICU and on- | Perform skin antisepsis | | Catheter days |
| | site neonatologist to | with povidone-iodine, | | Intervention: 30,587 |
| | join PQCNC CLABSI | CHG, or alcohol | | · · |
| | | Skin preparation agent | | Insertion compliance, %: |
| | Exclusion Criteria: NR | completely dry at time | | Baseline: 76 |
| | | of first skin puncture | | • Peaked: 93 |
| | | Procedure stopped if | | . cancar so |
| | | anyone notes sterility | | Insertion compliance, %, monthly (estimated |
| | | compromised | | from Figure): |
| | | · · | | October 2009: 76 |
| | | Catheter maintenance | | November 2009: 73 |
| | | checklist: | | December 2009: 87 |
| | | Volume of infant | | |
| | | feedings in mL/kg per | | January 2010: 92 |
| | | day | | February 2010: 90 |
| | | Central lines be | | March 2010: 93 |
| | | discontinued when | | April 2010: 92 |
| | | an infant's enteral | | May 2010: 88 |
| | | feedings reached | | June 2010: 80 |
| | | 120 mL/kg per day | | |
| | | Daily assessment of | | Maintenance compliance, %: |
| | | catheter need: | | Baseline: 32 |
| | | "Do we need the | | • Peaked: 56 |
| | | line today?" | | |
| | | "If there was no line | | Maintenance compliance, %, monthly |
| | | in place today, | | (estimated from Figure): |
| | | would we place | | October 2009: 32 |
| | | one?" | | November 2009: 40 |
| | | Dressing integrity and | | December 2009: 39 |
| | | site cleanliness | | |
| | | assessed (daily at | | January 2010: 38 |
| | | minimum) | | February 2010: 34 |
| | | Dressing and site care if | | March 2010: 34 |
| | | dressing change | | April 2010: 35 |
| | | performed | | May 2010: 56 |
| | | Site cleansed with | | June 2010: 46 |
| | | an appropriate | | |
| | | solution (povidone- | | Adverse events: NR |
| | | iodine, CHG, or | | |
| | | alcohol) | | |
| | | Cleansing solution | | |
| | | allowed to air-dry | | |
| | | completely | | |

| Study | Population and | Intervention/ Study | | |
|-------------|----------------|---|-------------|---------|
| Information | Setting | Groups | Definitions | Results |
| | | Use of a closed system: | | |
| | | closed system | | |
| | | maintained for infusion, | | |
| | | blood draws, and | | |
| | | medication | | |
| | | administration; closed | | |
| | | system is one in which | | |
| | | entries are made | | |
| | | through needleless | | |
| | | connectors or hubs that | | |
| | | have been disinfected | | |
| | | before use | | |
| | | For all catheter | | |
| | | entries/access | | |
| | | Scrub needleless | | |
| | | connector or hub | | |
| | | using friction with | | |
| | | alcohol or CHG for | | |
| | | ≥15 seconds | | |
| | | Allow surface of | | |
| | | connector or hub to | | |
| | | dry before entry | | |
| | | Staff wear clean | | |
| | | gloves when | | |
| | | accessing or | | |
| | | entering catheter (if | | |
| | | not using closed | | |
| | | system) | | |
| | | Control/Comparison: NA | | |
| | | Control/Companson. NA | | |
| | | Device/agent: NA | | |
| | | Device/agent. NA | | |
| | | Monitoring intervention: | | |
| | | Insertion and maintenance | | |
| | | compliance | | |
| | | | | |
| | | Standard preventive | | |
| | | measures: NR | | |

Table 58 Risk of Bias for Two Group Studies on Central Venous Catheter Insertion and Maintenance Bundles

| Author Year | All study groups derived from similar source/reference populations | Attrition not significantly different across study groups | Measure of exposure is valid | Measure of outcome is valid | Investigator blinded or were outcomes well- defined and objective to endpoint assessment | Potential confounders identified | Statistical adjustment for potential confounders done | Funding source(s) disclosed and no obvious conflict of interest | Overall Risk of Bias |
|------------------------------|--|--|------------------------------|-----------------------------|--|--|---|--|-------------------------|
| Balla 2018 ⁴⁷ | ✓ | ✓ | ✓ | ✓ | | | | ✓ | Moderate |
| Fisher 2013 ⁴⁵ | ✓ | | ✓ | ✓ | ✓ | | | ✓ | Moderate |
| Savage 2018 ⁴⁶ | ✓ | ✓ | ✓ | ✓ | √ | | | ✓ | Moderate |

C.13. Prophylactic Antimicrobial Administration

Key Question 13: In NICU patients requiring central venous catheters, what is the efficacy of prophylactic antimicrobials, compared with standard of care, to prevent CLABSI?

Table 59 Summary of Findings on Prophylactic Amoxicillin vs. No Prophylactic Amoxicillin to Prevent CLABSI

| Outcome | Findings | Quantity and Type of Evidence | GRADE of Evidence for Outcome and Limitations of the Evidence |
|--------------------------|--|---------------------------------------|---|
| Proven septicemia* | • One RCT ⁴⁸ found no difference was reported in proven septicemia (OR: 0.24; 95% CI: 0.01 -5.37 ; p = 0.37). | 1 RCT N=148 patients ⁴⁸ | Moderate • Imprecision: only one study |
| Suspected septicemia | • One RCT ⁴⁸ found no difference in suspected septicemia (OR: 0.47; 95% CI: 0.11 – 1.94; p = 0.29). | 1 RCT N=148 patients ⁴⁸ | Moderate • Imprecision: only one study |
| Thrombotic complications | • One RCT ⁴⁸ found thrombotic complications were reported in 9% of patients administered prophylactic amoxicillin, and 3% of the control group. | 1 RCT N=148 patients ⁴⁸ | Moderate • Imprecision: only one study |
| Amoxicillin resistance | One RCT⁴⁸ found one incidence of amoxicillin resistant Staphylococcus epidermidis in the control group. One RCT⁴⁸ found no decrease in amoxicillin susceptibility during the study period when compared with before the study period (47% vs. 42%), however susceptibility patterns after the study period were not reported. | 1 RCT N=148 patients ⁴⁸ | Moderate • Imprecision: only one study |

Table 60 Summary of Findings on Prophylactic Vancomycin vs. No Prophylactic Vancomycin to Prevent CLABSI

| Outcome | Findings | Quantity and Type of Evidence | GRADE of Evidence for Outcome and Limitations of the Evidence |
|--------------------------------|--|----------------------------------|---|
| CONS catheter- related sepsis* | • A reduction was seen in the incidence of CONS Catheter related sepsis (0/41 vs. 8/52 (15%); $p = 0.004$). | 1 RCT ⁴⁹ N=93 | Moderate • Imprecision: only one study |

| Outcome | Findings | Quantity and Type of Evidence | GRADE of Evidence for Outcome and Limitations of the Evidence |
|---|---|---|--|
| Laboratory confirmed BSI* | No difference was seen in the incidence of Laboratory Confirmed BSI in patients with peripheral CVCs for a period of prophylactic vancomycin compared with a period with no prophylaxis. (42/153 (27.4%) vs. 32/141 (22.7%); p = NS). This study reported an increase in the incidence of CONS BSI in patients with PCVCs when administered prophylactic vancomycin: 10/153 (6.5%) vs. 0/141 (0); P = 0.002). | 1 OBS ⁵⁰ N=294 | Very Low |
| Gram-positive infections | The use of prophylactic vancomycin for infants with central venous catheters was associated with reduced incidence of gram-positive infections (26/85 (31%) vs. 26/61 (43%); p<0.05). | 1 OBS ⁵¹ N=141 | Very Low ■ Study Quality: high risk of bias ■ Imprecision: only one study |
| Gram-negative infections | • One observational study ⁵¹ found the use of prophylactic vancomycin for infants with central venous catheters was associated with reduced incidence of gram-negative infections (19/85 (22%) vs. 21/61 (34%); p<0.05). | 1 OBS n=146 lines ⁵¹ | Very Low • Study Quality: high risk of bias • Imprecision: only one study |
| Total amount of vancomycin administered | One observational study⁵⁰ found that discontinuing prophylactic vancomycin resulted in fewer infants being exposed, but a larger total amount of vancomycin was administered for treatment of infection in the post-prophylactic period. | 1 OBS n=294 lines ⁵⁰ | Very Low ● Imprecision: only one study |
| Vancomycin Resistance | One RCT⁴⁹ reported no incidences of vancomycin resistance during the study, CONS susceptibility patterns did not change during study, and Vancomycin resistant strains of CONS were not detected during study. One observational study⁵¹ reported no incidences of vancomycin resistance were observed during the study period; however two years following the study, four cases of CONS resistance to vancomycin appeared. | 1 RCT N=93 lines ⁴⁹ 1 OBS n=146 lines ⁵¹ | Moderate • Imprecision: low number of events |

Table 61 Extracted Information on Prophylactic Antimicrobials

| Study | Population and Setting | Intervention/ Study Group | Definitions | Results |
|---------------------|--------------------------|--|--------------------------------------|-----------------------------------|
| Information | | | | |
| Author: | Number of Patients: | Intervention: | Outcome Definitions: | Primary Outcomes: |
| Harms ⁴⁸ | N=148 | n=75 | Proven Septicemia: Clinical signs | CLABSI: |
| | Number of lines: | Amoxicillin prophylaxis: 100mg/kg/ | (e.g., apnea, bradycardia, | Proven septicemia, n (%) |
| Year: | N = 148 | day in 3 doses, until catheter was | instability of temperature, | • Amoxicillin: 0/75 (0) |
| 1995 | | removed. | feeding problems, circulatory | • No Amoxicillin: 2/73 (2.7%) |
| | Setting: Neonatal ICU, | | changes, lethargy), suspect lab | Amoxicillin resistant: 1/2 (50%) |
| Study Design: | University Hospital | Control: | findings (CRP >0.6 mg/dl; I/T ratio | • OR: 0.24 (95% CI: 0.01 – 5.37); |
| RCT | | n=73 | >0.16), and cultures reveal | • p = 0.37 |
| | Location: Germany | No prophylactic antibiotics. | identical bacterial growth of the | · |
| Risk of Bias: | | | line tip and the blood. | Suspected septicemia, n (%): |
| Moderate | Dates: August 1990 - | Device/agent: Amoxicillin | | • Amoxicillin: 3/75 (4.0%) |
| | November 1992 | | Suspected septicemia: Clinical signs | • No Amoxicillin: 6/75 (8.2%) |
| | | Standard preventive measures: | and laboratory findings present | • OR: 0.47 (95% CI: 0.11 – 1.94); |
| | Inclusion Criteria: | Catheters inserted by a member | but no bacterial growth was | • p = 0.29 |
| | neonates with successful | of the medical staff using | identified in the culture of the | r |
| | central venous catheter | aseptic technique with infant in | blood specimen taken from the | Topic-specific outcomes: |
| | insertion. CVC insertion | the incubator. | peripheral vein. | -pp |

| Study | Population and Setting | Intervention/ Study Group | Definitions | Results |
|-------------|--|---|--|---|
| Information | .,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,, | | | |
| Information | was performed if peripheral venous access was difficult and the anticipated period of parenteral nutrition was longer than 10 days. Only initially inserted catheters were included in the analysis. Exclusion Criteria: NR | One unit of heparin added to each ml of the infusate. Blood products not administered through the catheter. Lines used to withdraw blood. Entire administration set, including all connectors, changed daily. Hub of the catheter and connecting pieces wrapped in sterile gauze. Catheters removed when no longer needed or when signs of serious infection, blockage or dislodgement occurred. Antibiotic therapy: Uniform regimen of abx therapy prescribed for all infants admitted to unit. Neonates with a history of infection, respiratory distress, clinical signs of infection, or suspect laboratory findings received combination intravenous amoxicillin and gentamicin therapy after blood culture specimens, tracheal aspirates, and skin swabs had been taken >90% of low birth weight or preterm neonates received antibiotic treatment initially. Treatment stopped after 48 - 72 hours if: cultures remained sterile, markers of inflammation were within the normal range, and no clinical signs of infection. In infants randomly assigned to receive prophylactic antibiotic treatment with amoxicillin, only the aminoglycoside was discontinued. | Mechanical complications: Clotting of catheter or dislodgement Sampling /Testing strategy: • A drop of fluid from the connecting hub was collected twice a week for bacteriologic examination. • Catheter tip removed cut off and placed immediately in nutrient broth for culture. Other notes: Every 10 infants, the study was evaluated. Decision to stop or continue depended on indication of superiority of amoxicillin treatment or if superiority could not be proved. | Duration of catheterization, median, days (25 th to 75 th percentiles): • Amoxicillin: 15 (10-23) • No amoxicillin: 15 (12-25) Adverse events Antibiotic susceptibility of all isolated microorganisms (in vitro): • During study period: 47% • Before study period: 42% Thrombotic complications, n (%): • Amoxicillin: 7/75 (9.3%) • No amoxicillin: 2/73 (2.7%) • p = NR . Mechanical complications, n (%): • Amoxicillin: 3/75 (4.0%) • No amoxicillin: 4/73 (5.5%) • p = NR Thrombocytopenia <150, n (%): • Amoxicillin: 7/75 (9.3%) • No amoxicillin: 9/73 (12.3%) • p = NR CRP >0.6 mg/dl, n (%): • Amoxicillin: 8/75 (10.6%) • No amoxicillin: 10/73 (13.6%) • p = NR I/T ratio >0.16, n (%): • Amoxicillin: 14/75 (18.6%) • No amoxicillin: 16/73 (21.9%) • p = NR Additional antibiotics, n (%): • Amoxicillin: 20/75 (26.7%) • No amoxicillin: 18/73 (24.7%) • p = NR |

| Study Information | Population and Setting | Intervention/ Study Group | Definitions | Results |
|------------------------|------------------------------|---|--|---|
| Information | | . If infants had sings of | | |
| | | If infants had signs of page against infantion, they | | |
| | | nosocomial infection, they received cefotaxime or | | |
| | | ceftazidime and netilmicin, | | |
| | | | | |
| | | amikacin, or tobramycin. | | |
| | | Other abx (e.g., vancomycin) | | |
| | | administered according to the | | |
| | | susceptibility of the isolated | | |
| Author: | Number of patients: | organism. Intervention: n=35 patients; | Outcome Definitions: | Primary Outcomes: |
| Spafford ⁴⁹ | N = 70 | n=41 catheters | Catheter related sepsis: When the | CONS Catheter related sepsis, No. of catheters, n (%): |
| Sparioru | Number of lines: | | culture of the CVC specimen | • Vancomycin: 0/41 (0) |
| Year: | N = 93 | • TPN with 25 μg/ml Vancomycin | contained at least 10 times the | • No vancomycin: 8/52 (15%) |
| 1994 | N = 33 | Control: n=35 patients; | concentration of the same | , , , , , |
| 1994 | Setting: Neonatal ICU, | N=52 catheters | pathogen isolated from the | • p = 0.004 |
| Study Design: | Regional Hospital | • TPN only | peripheral sample. | Non-CONS Catheter related sepsis, No. of catheters, n |
| Prospective, | Regional Hospital | TPN Offity | Infants examined for sepsis | (%): |
| double blind | Location: USA | Device/agent: Vancomycin | when they had temperature | • Vancomycin: 1/41 (2.4%) |
| RCT | Location: 65/ | Device/agent. vancomycm | instability, increased oxygen or | • No vancomycin: 5/52 (9.6%) |
| | Dates: April 1991- June | Standard preventive measures: | ventilator requirements, | • p = NR |
| Risk of Bias: | 1992 | Catheters placed under sterile | increased number or severity of | Tania anasifia automasa |
| Low | 1551 | conditions. | episodes of apnea or | Topic-specific outcomes: |
| | Inclusion Criteria: All | Catheters were inserted only | bradycardia, feeding | Duration of catheterization, mean days (±SE): |
| | infants admitted to the | after a negative blood culture | intolerance, lethargy, or blood | • Vancomycin: 18.7 (±5.4) |
| | NICU in whom a CVC was | finding had been obtained, and | pressure instability. If sepsis | No vancomycin: 17.3 (±2.5) |
| | inserted. (general care for | there was no evidence of an | suspected, blood specimens | • p = NS |
| | infants weighing <1000g | acute infection | obtained from peripheral vein | Adverse events |
| | included insertion of a CVC | Insertion site covered with a | and through CVC | Antibiotic resistance: |
| | on day 3 or 4 to improve | clear bio-occlusive dressing that | Sampling /Testing strategy: | CONS susceptibility patterns: did not change during study |
| | overall nutrition.) | was changed only if necessary. | If sepsis was suspected, blood | Vancomycin resistant strains of CONS: not detected |
| | | All infants given empiric | culture specimens obtained | during study |
| | Exclusion Criteria: Broviac, | treatment with ampicillin and | from a peripheral vein and | during study |
| | Hickman or umbilical | gentamicin at birth. | drawn through the CVC were | BUN, mmol/L (mg/dl): |
| | venous catheters were not | These antimicrobial agents were | obtained. | • Vancomycin: 6.5 (18.2) |
| | included as study catheters | continued until culture results | On removal, catheters were | • No vancomycin: 6.5 (18.2) |
| | and were not used in | were confirmed negative at 48 | sent to the microbiology | • p = NS |
| | conjunction with a CVC. | hours after birth. | laboratory for culture of | - P 110 |
| | Infants with renal | • TPN solution infused over 24h | catheter specimens to | Creatinine, μmol/L (mg/dl): |
| | dysfunction. | Ampicillin and gentamicin used | determine colonization. | • Vancomycin: 80 (0.9) |
| | | during periods of suspected | Concentrations of blood urea | • No vancomycin: 88 (1.0) |
| | | sepsis, for 48 hours pending | nitrogen were measured each | • p = NR (noted not different) |
| | | results of cultures. If a positive | week to assess renal function. | - p - W (noted not different) |
| | | culture, then appropriate | Vancomycin concentrations | |

| Study Information | Population and Setting | Intervention/ Study Group | Definitions | Results |
|---|---|---|---|---|
| | | antimicrobial therapy continued for 10 days. • Vancomycin administered only for culture-proven positive infections | measured weekly. Brain-stem auditory evoked responses were obtained before discharge to determine possible vancomycin-induced ototoxic effects. Other notes: Majority of catheters inserted at 48-96 h of age to provide concentrated TPN solution. | Mortality, n: • Vancomycin: 5/35 (sepsis: 0) • No vancomycin: 9/35 (sepsis: 4/9, none attributable to CVC) • p = NR |
| Author: Elhassan ⁵⁰ Year: 2004 Study Design: Uncontrolled before after (Retrospective Cohort) Risk of Bias: High | Number of patients: N = 294 Number of lines: N = 294 Setting: Neonatal ICU, Tertiary Care Hospital Location: USA Dates: June 1, 1997 – September 31, 2000: • Period I: June 1, 1997 - December 31, 1998 • Period II: April 1, 1999 - September 31, 2000 Inclusion Criteria: Neonates admitted to the NICU during the study periods and had a PCVC inserted during their stay. Infants with UVC placed before PCVC. Exclusion Criteria: Infants with surgically placed catheters (Broviac or Hickman) or femoral. | Patient Groups: Period I: n= 153 patients; n=193 catheters Prophylactic Vancomycin in Hyperalimentation solutions (HAL) Period II: n=141 patients; n=178 catheters No Prophylactic Vancomycin, Device/agent: Vancomycin Standard preventive measures: PCVCs inserted in the NICU percutaneously through a needle or under direct visualization of the vein through a cutdown technique. No change in catheter management technique between study periods | Outcome Definitions: Nosocomial laboratory confirmed blood stream infections (LC-BSI): if a (+) blood culture was collected beyond 3 days of age and the patients satisfied Criterion I, or IIa or IIb and positive lab results are not related to an infection at another site. • Criterion I- Patient has a recognized pathogen cultured from one or more blood cultures, and the organisms cultured from blood are not related to an infection at another site. • Criterion II- Patient age <1 year has at least one of the following signs or symptoms: fever >100.4°F, hypothermia <98.6°F, apnea or bradycardia and at least one of the following: • Criterion IIa- common skin contaminants cultured from two or more blood cultures drawn on separate occasions; • Criterion IIb- common skin contaminants cultured from at least one blood culture from a patient with an intravenous | Primary Outcomes: LC-BSI, total no. of positive blood cultures; n (%): Period I (proph): 52/153 (34.0%) Period II (no proph): 64/141 (45.3%) p = 0.0457 Group A (with PCVC), LC-BSI, total no. of positive blood cultures; n (%): Period I (proph): 42/153 (27.4%) Period II (no proph): 32/141 (22.7%) p = NS Group B (no PCVC), LC-BSI, total no. of positive blood cultures; n (%): Period I (proph): 10/153 (6.5%) Period II (no proph): 26/141 (18.4%) p = 0.0019 Topic-specific outcomes: Duration of catheterization, mean days (SD): Period I (proph): 22.1 (±19.2) Period II (no proph): 20.8 (±15.4) p = NS Patients given Prophylactic Vancomycin, n: Period I (proph): 151/153 Period II (no proph): 0/141 p = NR Amount of vancomycin administered, mean (g): Period I (proph): 5.85 Period II (no proph): 0 p = NR |

| Study | Population and Setting | Intervention/ Study Group | Definitions | Results |
|-------------|------------------------|---------------------------|---|---|
| Information | | | institutes appropriate antimicrobial therapy; and signs and symptoms with positive laboratory results are not related to an infection at another site. Group A: With PCVC in place Group B: Without PCVC in place. Cultures collected before PCVC insertion or up to 7 days after PCVC removal Effect of continuous vancomycin prophylaxis evaluated through HAL on: 1) total count and longevity of PCVCs and 2) the total vancomycin exposure in the two periods. Sampling /Testing strategy: Blood cultures. Other notes: None | Total number and rate of patients who received vancomycin treatment, n (%): Period I (proph): 29/153 (18.9%) Period II (no proph): 43/141 (30.4%) p = 0.0215 Vancomycin treatment for Proven LC-BSI, n (%): Period I (proph): 14/153 (9.1%) Period II (no proph): 24/141 (17.0%) p = 0.0025 Amount of vancomycin administered, for Proven LC-BSI mean (g) Period I (proph): 2.72 Period II (no proph): 10.0 p = NS Vancomycin treatment for Suspected Infection n, (%) Period I (proph): 15/153 (9.8%) Period II (no proph): 19/141 (13.5%) p = NS Amount of vancomycin administered for Suspected Infection, n (g) Period I (proph): 2.35 Period II (no proph): 4.29 p = NS Adverse events LC-BSI by organism, no. of positive blood cultures; n (%): Coagulase-negative Staphylococcus, n (%): Period I (proph): 19/153 (12.4%) Period II (no proph): 31/141 (21.9%) p = 0.0291 Group A, n (%): Period II (no proph): 25/141 (17.7%) p = NS Group B, n (%): Period II (no proph): 3/153 (2.0%) Period II (no proph): 6/141 (4.2%) p = NS Other gram-positive organisms Period II (no proph): 14/141 (9.9%) p = NS |

| Study | Population and Setting | Intervention/ Study Group | Definitions | Results |
|--------------------------------|---|---|---|---|
| Information | ropulation and Setting | intervention, study Group | Demittoris | Group A, n (%): Period I (proph): 7/153 (4.5%) Period II (no proph): 8/141 (5.7%) p = NS Group B, n (%): Period I (proph): 0/153 (0) Period II (no proph): 6/141 (4.2%) p = 0.0099 Gram-negative organisms Period I (proph): 15/153 (9.8%) Period II (no proph): 9/141 (6.4%) p = NS Group A, n (%): Period I (proph): 10/153 (6.5%) Period II (no proph): 0/141 (0) p = 0.002 Group B, n (%): Period II (no proph): 5/153 (3.3%) Period II (no proph): 9/141 (6.4%) p = NS Fungal organisms Period I (proph): 11/153 (7.2%) Period II (no proph): 10/141 (7.1%) p = NS Group A, n (%): Period I (proph): 9/153 (5.9%) Period II (no proph): 5/141 (3.5%) p = NS Group B, n (%): |
| | | | | Period I (proph): 2/153 (1.3%) Period II (no proph): 5/141 (3.5%) p = NS |
| Author: Ocete ⁵¹ | Number of patients: N = 146 | Intervention: n= 85 Prophylactic Vancomycin at 25 | Outcome Definitions: Infection: with presence of at least | Primary Outcomes: Infections, n [numerator calculated by CDC] (%): |
| | No differences between | μg/mL through catheter | two clinical symptoms (bad | Negative coagulase staphylococci (NCS) |
| Year: 1998 | the two groups in terms of gestational | Control: n= 61 | perfusion, apnea, respiratory distress, digestive, neurological, or | Vancomycin: 19/85 (22%)No vancomycin: 21/61 (34%) |
| | age, weight, risk factors | No Prophylactic Vancomycin | urinary disorders) in the absence of | • p < 0.05 |
| Study Design: | on admittance or | | any other evidence cause of the | Gram positive |
| Non- | duration of assisted respiration. | Device/agent: Vancomycin | clinical alteration. | Vancomycin: 26/85 (31%) No vancomycin: 26/61 (43%) |

| Study | Population and Setting | Intervention/ Study Group | Definitions | Results |
|---|---|---|--|---|
| Randomized Control Study Risk of Bias: High | Intervention group contained a higher number of newborns with assisted respiration (p<0.01). Number of lines: N = 146 Setting: Neonatal ICU, university hospital Location: Spain Dates: Control: September 10, 1993 - September 9, 1994 Intervention: September 10, 1994 - September 9, 1995 Inclusion Criteria: Newborns admitted to the NICU requiring central catheters (umbilical artery, umbilical vein and/or silastic) during the study periods for both groups. | Standard preventive measures: Umbilical and silicone catheters inserted using sterile technique with povidone iodine applied to all connections. Umbilical catheters fitted by doctor and Silicone catheters fitted by nurse. | Sampling /Testing strategy: Central and peripheral cultures were performed. Other notes: None | p < 0.05 Gram negative Vancomycin: 19/85 (22%) No vancomycin: 20/61 (33%) p = NS Fungus Vancomycin: 6/85 (7%) No vancomycin: 6/61 (10%) p = NS Topic-specific outcomes: Duration of catheterization, mean days (SD): Vancomycin: 9.20 (±9.15) No vancomycin: 9.36 (±13.35) p = NS Adverse events Antibiotic resistance: No resistance to vancomycin observed during the study period. Two years following the study, four cases of NCS resistance to vancomycin appeared. |
| | Exclusion Criteria: NR | | | |

Table 62 Risk of Bias for Randomized Controlled Trials on Prophylactic Antimicrobials

| Author Year | Described as randomized | Randomization appropriately performed | Described as double- blind | Outcome assessor blinded | Study participant blinded | Investigator blinded | Attrition described | Attrition smaller than 10-15% of assigned patients | Attrition appropriately analyzed | Funding source(s) disclosed and no obvious conflict of interest | Overall Risk of Bias |
|--------------------------------|-------------------------------|---------------------------------------|----------------------------------|--------------------------------|---------------------------------|-------------------------|------------------------|---|--|---|-------------------------|
| Harms 1995 ⁴⁸ | ✓ | ✓ | | | | | ✓ | ✓ | ✓ | | Moderate |
| Spafford 1994 ⁴⁹ | ✓ | ✓ | ✓ | ✓ | ✓ | ✓ | ✓ | ✓ | ✓ | | Low |

Table 63 Risk of Bias for Two Group Studies on Prophylactic Antimicrobials

| Author Year | Were patients | For non-randomized trials, did the study employ any other methods to enhance group comparability such as matching, stratification, or statistical methods to adjust for baseline differences? | study groups have similar levels of performance on the | Did the study enroll all suitable patients or consecutive | Was the comparison of | Were the two groups treated/ evaluated concurrently? | • | Was the funding for this study derived from a source that would not benefit financially from results in a particular direction? | Risk of Bias |
|-----------------------------|---------------|---|--|--|-----------------------|---|---|---|--------------|
| Elhassan | Вісирої | umerenees | √ | <i>√</i> | piamica | ochoun chu, | | un cononi | High |
| 2004 ⁵⁰ | | | • | ŕ | | | | | riigii |
| Ocete 1998 ⁵¹ | | | ✓ | ✓ | ✓ | | | | High |

C.14. Prophylactic Anticoagulant Administration

Key Question 14: In NICU patients requiring central venous catheters, what is the efficacy of prophylactic anticoagulant infusions, compared with standard of care, to prevent CLABSI?

Table 64 Summary of Findings on Prophylactic Heparin + TPN or dextrose vs. TPN or dextrose to Prevent CLABSI

| Outcome | Findings | Quantity and Type of Evidence | GRADE of Evidence for Outcome and Limitations of the Evidence |
|---|---|---|---|
| Catheter-related sepsis (CRS) or definite CRS* | Four RCTs ⁵²⁻⁵⁵ found no difference in the incidence of catheter-related sepsis or definite CRS when comparing the use of prophylactic heparin with no heparin. | 4 RCT ⁵²⁻⁵⁵ N=210 patients N=66 patients N=201 patients N=239 patients | High |
| Definite or probable CRS* | • One RCT study found no difference in the incidence definite or probable CRS when comparing the use of heparin with no heparin. [9/102 vs. 16/108; RR: 0.60 (95% CI: 0.28 – 1.26); p = 0.18]. | 1 RCT ⁵² N=210 patients | Moderate • Imprecision: only one study |
| Septicemia* | • One RCT study found no difference in the incidence of septicemia when comparing the use of heparin with no heparin. [7/35 (20.0%) vs. 9/31 (29.0%); RR: 0.7 (95% CI: 0.3-1.6); p = NR]. | 1 RCT ⁵⁵ N=239 patients | Moderate • Imprecision: only one study |
| Occlusion | Two RCT studies^{52, 53} found no difference in the incidence of occlusion with the use of heparin compared with no heparin [5/102 vs. 3/108; RR: 1.76 (95% CI: 0.48-6.56); p = 0.42] & [5/35 (14.3%) vs. 7/31 (22.6%); RR: 0.6 (95% CI: 0.2-1.8); p = NR]. Two RCT studies^{54, 55} found heparin was associated with significant reduction in occlusion (23/118 (19.5%) vs. No heparin: 55/121 (45.5%); RR: 3.44 (95% CI: 1.92-6.44); p<0.05 (=0.0001)] & [6/100 vs. 31/101; RR: 0.20 (95% CI: 0.09-0.42); p<0.05 (=0.001)]. | 4 RCT ⁵²⁻⁵⁵ N=210 patients N=66 patients N=239 patients N=201 patients | Moderate ■ Consistency: inconsistent results |
| Intraventricular hemorrhage | Three RCT studies ⁵²⁻⁵⁴ reported no difference in the incidence of intraventricular hemorrhage with the implementation of prophylactic anticoagulant. | 3 RCT ⁵²⁻⁵⁴ N=210 patients N=66 patients N=201 patients | High |

Table 65 Extracted Information on Anticoagulant Infusion

| Study Information | Population and Setting | Intervention | Definitions | Results |
|-----------------------------|---------------------------------|--|---|--|
| Author: Birch ⁵² | Number of patients: | Intervention: n=102 | Outcome Definitions: | Primary Outcomes: |
| | N = 210 | Heparin plus TPN | Catheter related sepsis (CRS): A | Definite catheter related sepsis, n: |
| Year: 2010 | Number of lines: | | positive blood culture growing CONS, | • Heparin: 3/102 |
| | N = 210 | Control: | Staphylococcus aureus, | • No heparin: 10/108 |
| Study Design: | | n=108 | Acinetobacter species or Candida. | • RR: 0.32 (95% CI: 0.1-1.03) |
| Prospective | Setting: Tertiary Neonatal ICU | TPN without heparin | | • p = 0.06 |
| double blind RCT | , | · | Definite CRS: Two positive blood | ν μ – 0.00 |
| | Location: New Zealand | Device/agent: Heparin | cultures with the same organism | Rates of definite catheter related sepsis/1000 days |
| Risk of Bias: Low | | | taken from two separate sites within | catheter in situ, n: |
| | Dates: March 2004-October | Monitoring intervention: | 72 hours of each other. | • Heparin: 2.3 |
| | 2007 | | | No heparin: 6.8 |
| | | Standard preventive | Probable CRS: Single positive blood | • RR: 0.34 (95% CI: 0.09-1.24) |
| | Inclusion Criteria: Infants | measures: | culture and a peak C-reactive protein | • p = 0.09 |
| | requiring a long line for TPN | Long lines were | level greater than 9 mg/l recorded | Ψ p = 0.09 |
| | as judged by the clinical team | inserted according to | from 24 h before to 72 h after the | Probable catheter related sepsis, n: |
| | | current unit practice | positive culture was drawn. | • Heparin: 6/102 |
| | Exclusion Criteria: Any | using an aseptic | | • No heparin: 6/108 |
| | previous long line successfully | technique and all lines | Possible CRS: Single positive blood | • RR: 1.06 (95% Cl: 0.37-3.03) |
| | inserted and utilized | were secured using | culture without elevation of C- | |
| | | medical adhesive and | reactive protein. | • p = 0.92 |
| | | covered with non- | | Descible eathers related consist no |
| | | adhesive dressing. | Bacteremia with organisms not | Possible catheter related sepsis, n: |
| | | | commonly associated with line | • Heparin: 6/102 |
| | | Choice of catheter was | sepsis: a single positive blood culture | • No heparin: 13/108 |
| | | determined by the | with the following organisms: | • RR: 0.49 (95% CI: 0.2-1.19) |
| | | inserting physician. | streptococcal species, Gram-negative | • p = 0.12 |
| | | Following insertion, | organisms and enterococci. Two or | A 606/16:3 |
| | | the lines were either | more blood cultures positive for the | Any CRS (definite, probable, possible), n: |
| | | attached directly to a | same organism and less than 7 days | • Heparin: 15/102 |
| | | bag of TPN or to an | apart were considered to be the | No heparin: 28/108 |
| | | infusion of normal | same single bacteremia episode. | • RR: 0.57 (95% CI: 0.32-0.98) |
| | | saline while waiting | | • p<0.05 (=0.04) |
| | | for the confirmation | Positive blood culture: any blood | D. Conclusion I. Conclusion |
| | | of the position of the | culture growing one or more | Rate: any episodes of CRS/1000 days catheter in situ, n: |
| | | line. | organism drawn from insertion of | • Heparin: 12.3 |
| | | | the long line to 24 hours after the | No heparin: 20.3 |
| | | | line was removed. | • RR: 0.61 (95% CI: 0.33-1.11) |
| | | | | • p = 0.10 |
| | | | Intraventricular hemorrhage (IVH) | |
| | | | progression: an increase on either | Definite or probable CRS, n: |
| | | | side from grade 0–2 to grade 3–4 | |

| Study Information | Population and Setting | Intervention | Definitions | Results |
|-------------------|------------------------|--------------|--|---|
| Study Information | Population and Setting | Intervention | between the 'worst initial IVH' and the 'worst post-trial IVH'. Sampling /Testing strategy: Blood cultures Other notes: None | Results ● Heparin: 9/102 ● No heparin: 16/108 ● RR: 0.60 (95% CI: 0.28 – 1.26) ● p = 0.18 Bacteremia with organisms not commonly associated with line sepsis, episodes: ● Heparin: 1 ● No heparin: 0 ● p = NR Topic-specific outcomes: Duration of catheter patency, mean days (SD): ● Heparin: 12.9 (±9.8) ● No heparin: 13.7 (±12.4) ● p = 0.93 Adverse events: Occlusion, n: ● Heparin: 5/102 ● No heparin: 3/108 ● RR: 1.76 (95% CI: 0.48-6.56) ● p = 0.42 Extravasation, n: ● Heparin: 4/102 ● No heparin: 8/108 ● RR: 0.53 (95% CI: 0.17-1.6) ● p = 0.28 IVH Progression, n: ● Heparin: 2/102 ● No heparin: 7/108 ● RR: 0.3 (95% CI: 0.07 - 1.24) ● p = 0.11 |
| | | | | Non-catheter-related sepsis, n: • Heparin: 1/102 • No heparin: 0/108 • p = NR Mortality, n: • Heparin: 0/102 |

| Study Information | Population and Setting | Intervention | Definitions | Results |
|--|--|--|--|--|
| | | | | No heparin: 1/108 p = NR Bleeding diatheses: None observed |
| | | | | Thrombocytopenia: None observed |
| Author: Uslu ⁵⁵ | Number of patients: N = 239 | Intervention group: n=118 | Outcome Definitions: Catheter related sepsis: Clinical signs of | Primary Outcomes: Catheter related sepsis, n (%): |
| Year: 2010 | Number of lines: N = 239 | Heparin plus TPN | sepsis was associated with a positive peripheral blood culture and positive | Heparin: 2/118 (1.7)No heparin: 4/121 (3.3) |
| Study design: Prospective double blind RCT | Setting: Neonatal ICU | Control group: n=121 TPN without heparin | catheter culture of the same organism. Duration of catheter: Number of days | • p = 0.68 |
| Risk of Bias: Low | Location: Turkey Dates: February 1, 2007- October 31, 2008 | Device/agent: Heparin Monitoring intervention: | between insertion and removal. Catheter removal: signs of local or systemic infection, phlebitis, | Septicemia, n (%): • Heparin: 5/118 (4.2) • No heparin: 4/121 (3.3) • p = 0.74 |
| | Inclusion Criteria: All neonates admitted to the NICU who had required a peripherally inserted percutaneous central venous catheter (PCVC) as determined by the attending neonatologist. Exclusion Criteria: Neonates | Standard preventive measures: • Catheters were placed by using a sterile technique. Catheter type and place of insertion were determined by the physician's choice. • Catheters were | extravasation, blockage, breakage and leakage of catheter, accidental removal, death, and if neonate reached close to full enteral feeds Catheter occlusion: the inability of infusing fluids through the catheter due to blockage Thrombosis: a thrombus along the catheter line detected by inspection | Topic-specific outcomes: Duration of catheter patency, days: • Heparin: 12.4 (±4.5) • No heparin: 9.7 (±4.0) • p < 0.05 (=0.0001) Adverse events: Occlusion, n (%): • Heparin: 23/118 (19.5) |
| | with bleeding tendencies, grade 3 to 4 intraventricular hemorrhage, recent suspected or confirmed sepsis (within 48 h of initiation of antibiotic therapy), thrombocytopenia (<100,000 mm ⁻³), disseminated intravascular coagulation, arrhythmia and congonital | stabilized and secured with a transparent medical film dressing, which was not changed unless it became polluted or slack. | after removal of the catheter Phlebitis: inspection as swelling, hyperemia and change in skin color associated with an inflamed vein Sampling /Testing strategy: Bacterial cultures were obtained from catheters and flushing solutions. In case of | No heparin: 55/121 (45.5) RR: 3.44 (95% CI: 1.92-6.44) p < 0.05 (=0.0001) Thrombosis, n (%): Heparin: 2/118 (1.7) No heparin: 5/121 (4.1) p = 0.25 Phlebitis, n (%): |
| | arrhythmia, and congenital malformations. Additionally, patients with uncertain viability | | suspicion of septicemia, blood culture was obtained. Other notes: None | Heparin: 10/118 (8.4) No heparin: 10/121 (8.3) p = 0.12 |

| Study Information | Population and Setting | Intervention | Definitions | Results |
|----------------------------|--------------------------------|---------------------|----------------------|---|
| | (determined by | | | Thrombocytopenia, n: |
| | neonatologist), need for use | | | • Heparin: 2/118 |
| | of heparin (umbilical arterial | | | • No heparin: 1/121 |
| | catheter), and a prolonged | | | • p = NR |
| | activated partial | | | · |
| | thromboplastin time (aPTT) | | | aPTT >100s, n: |
| | (>74 s for preterm infants and | | | • Heparin: 1/118 |
| | >51 s for term infants) | | | No heparin: 0/121 |
| | | | | • p = NR |
| | | | | Bleeding tendencies, n: |
| | | | | • Heparin: 1/118 |
| | | | | • No heparin: 1/121 |
| | | | | • p = NR |
| | | | | • |
| | | | | Intracranial hemorrhage, n (%): |
| | | | | Heparin: 19/118 (16.1) |
| | | | | No heparin: 21/121 (17.4) |
| | | | | • p = 0.93 |
| | | | | Intracranial hemorrhage after PCVC removal, n (%): |
| | | | | • Heparin: 21/118 (17.8) |
| | | | | • No heparin: 23/121 (19.0) |
| | | | | • p = 0.80 |
| | | | | p 0.00 |
| | | | | Arrythmia after PCVC removal, n (%): |
| | | | | • Heparin: 1/118 (0.8) |
| | | | | • No heparin: 1/121 (0.8) |
| | | | | • p = 0.80 |
| | | | | |
| | | | | Mortality, n (%): |
| | | | | • Heparin: 6/118 (5.1) |
| | | | | • No heparin: 6/121 (4.8) |
| | | | | • p = 0.79 |
| | | | | Other (e.g., breakage, leakage, accidental withdrawal), n |
| | | | | (%): |
| | | | | • Heparin: 3/118 (2.5) |
| | | | | • No heparin: 4/121 (3.2) |
| | | | | • p = 1 |
| Author: Shah ⁵⁴ | Number of patients: | Intervention: n=100 | Outcome Definitions: | Primary Outcomes: |
| | N = 201 | | | Catheter related sepsis, n: |

| Study Information | Population and Setting | Intervention | Definitions | Results |
|---|---|--|--|---|
| Year: 2007 Study Design: Prospective double blind RCT Risk of Bias: Low | Number of lines: N = 201 Setting: Four tertiary care Neonatal ICUs Location: Canada Dates: November 2002- November 2005 Inclusion Criteria: All neonates requiring peripherally placed percutaneous central venous catheters (PCVC) access as judged by the clinical team Exclusion Criteria: Neonates who had grade ¾ intraventricular hemorrhage, recent onset of presumed or confirmed sepsis (within 48 hours of initiation of antimicrobial therapy), bleeding diathesis, disseminated intravascular coagulation, thrombocytopenia, arrhythmia, or preexisting liver disease. | Heparin: 10% or 5% dextrose with heparin Control: n=101 No heparin: 10% or 5% dextrose Device/agent: Heparin Monitoring intervention: Standard preventive measures: • All PCVCs were placed by using sterile technique as per similar standards in each NICU. • Catheters were flushed by normal saline before insertion, and the extension tubing was connected to the PCVC hub. • Catheters were secured by transparent occlusive dressing that was not changed unless it was soiled or loose | Catheter related sepsis: Symptoms and signs suggestive of sepsis with a positive blood culture obtained from catheter fluid and a normally sterile site (blood urine, or cerebrospinal fluid) for the same organism. Catheter occlusion: the inability to infuse fluid Duration of catheter use: time between insertion and removal (elective or because of complications) of the catheter in hours. Thrombosis: the detection of a thrombus along the catheter path Sampling /Testing strategy: NR Other notes: None | Heparin: 5/100 No heparin: 2/101 p = 0.243 Suspected catheter-related sepsis, n: Heparin: 5/100 No heparin: 4/101 OR: 1.28 (95% CI: 0.33-4.90) p = 0.722 Topic-specific outcomes: Duration of catheter patency, mean hours (SD): Heparin: 267 (±196) No heparin: 233 (±194) p = 0.220 Duration of catheter patency, median (range): Heparin218 (6-1095) heparin No heparin: 188 (3-1176) p = NR Duration of catheter usability, n: p < 0.05; Hazard ratio: 0.53 (95% CI: 0.35-0.81) Adverse events: Reasons for non-elective catheter removal Occlusion, n: Heparin: 6/100 No heparin: 31/101 RR: 0.20 (95% CI: 0.09-0.42) p < 0.05 (=0.001) Non occlusive thrombosis, n: Heparin:18/100 No heparin: 18/101 p = NR Intraventricular hemorrhage: None observed HIT thrombocytopenia, n: Heparin: 1/100 No heparin: 0/101 p = NR Bleeding: |

| Study Information | Population and Setting | Intervention | Definitions | Results |
|------------------------------|--|---|--|---|
| | | | | None observed |
| | | | | Leakage, n: |
| | | | | • Heparin: 6/100 |
| | | | | No heparin: 2/101 |
| | | | | • p = 0.145 |
| | | | | Extravasation, n: |
| | | | | • Heparin: 8/100 |
| | | | | • No heparin: 14/101 |
| | | | | • p = 0.183 |
| | | | | Other reasons for non-elective catheter removal, n: |
| | | | | • Heparin: 7/100 |
| | | | | No heparin: 6/101 |
| | | | | • p = 0.760 |
| Author: Kamala ⁵³ | Number of patients: | Intervention group: n=35 | Outcome Definitions: | Primary Outcomes: |
| | N = 66 | Heparin plus TPN | Catheter related sepsis: Present in | Catheter related sepsis, n (%): |
| Year: 2002 | Number of lines: | | neonates manifesting clinical signs of | Heparin:1/35 (2.9) |
| Study Design: | N = 66 | Control group: n=31 | sepsis associated with a positive | • No heparin: 1/31 (3.2) |
| Prospective | | TPN no heparin | catheter-tip culture and a positive | • RR: 0.9 (95% CI: 0.06-13.6) |
| double-blind RCT | Setting: Neonatal ICU | | peripheral blood culture of the same | • p = NR |
| | | Device/agent: Heparin | bacterial organism. | |
| Risk of Bias: Low | Location: Malaysia | | | Septicemia, n (%): |
| | | Monitoring intervention: | Septicemia: Diagnosed when infants | Heparin: 7/35 (20.0) |
| | Dates: August 1,1999-August | | developed clinical signs of sepsis | • No heparin: 9/31 (29.0) |
| | 31, 2000 | Standard preventive | associated with a positive blood | • RR: 0.7 (95% CI: 0.3-1.6) |
| | | measures: | culture, irrespective of the catheter tip | • p = NR |
| | Inclusion Criteria: All | The TPN fluids used in | culture result. | |
| | neonates admitted to the | both groups of infants | | Topic-specific outcomes: |
| | NICU who had Peripherally or | were prepared under | Duration of catheter patency: the | Duration of PICC in situ, mean days (SD): |
| | percutaneously inserted | sterile conditions by | number of days for which the PICC | • Heparin: 10.8 (±6.7) |
| | central venous catheters | the pharmacist. | remained functioning in-situ, and upon | • No heparin: 9.3 (5.1) |
| | (PICCs) inserted subsequently | Catheters were | removal there was no evidence of | • 95% CI difference between means: -4.4-1.4 |
| | for the purpose of receiving TPN. | inserted | blockage. | • p = NR |
| | IFIV. | percutaneously from a | Hyperhiliruhinemia: Diagnosed as | |
| | Exclusion Criteria: Neonates | sterile protective | Hyperbilirubinemia: Diagnosed as being present when any infant's serum | Adverse events |
| | with clinical evidence of | conduit through either a 21 or 19 gauge | bilirubin level rose higher than normal | Blocked catheter/ Occlusion, n (%): |
| | bleeding tendencies, severe | winged needle. | Sim abin level 103e mgner than normal | • Heparin: 5/35 (14.3) |
| | IVH of grade 3 or 4; platelet | williged fieldlie. | Sampling /Testing strategy: Specimens | • No heparin: 7/31 (22.6) |
| | counts <100 x 10 ⁹ 1 ⁻¹ and/or | | of blood was collected from each | • RR: 0.6 (95% CI: 0.2-1.8) |
| | prolonged activated partial | | infant for measurement of bilirubin, | • p = NR |
| | thromboplastin time (APTT | | triglyceride, APTT and platelet count | |
| | more than 51 sec for term | | before insertion of catheter and again | Intraventricular hemorrhage, n (%): |
| | infants of gestation ≥37 | | on days 4 and 8 with PICC in situ, or on | Heparin: 4/23 (17.4) |

| Study Information | Population and Setting | Intervention | Definitions | Results |
|-------------------|--|--------------|--|---|
| | weeks, or more than 74 sec for preterm infants of gestation <37 weeks. | | removal of the PICC if the catheter was to be removed before day 4. Catheter blockage was diagnosed when unable to infuse TPN fluid readily through the catheter while in situ and detection of clots in the PICC after removal from the infants. If clot was detected upon removal, the catheter tip and aseptically collected solution were sent for bacterial culture. A specimen of blood for bacterial culture was obtained from the peripheral vein of an infant whenever attending doctor suspected septicemia. Cranial ultrasonography was carried out before, 1 week after commencement and upon completion of TPN. Other notes: None | No heparin: 4/20 (20.0) RR: 0.9 (95% CI: 0.3-3.00) p = NR Peak serum bilirubin level, mean μmol 1⁻¹ (SD): Heparin: 199 (±65) No heparin: 230 (±71) 95% CI difference between means: -1.4-63.8 p = NR Peak serum triglyceride level, mean mmol 1⁻¹ (SD): Heparin: 2.3 (±1.5) No heparin: 1.9 (±1.4) 95% CI difference between means: -1.2-0.3 p = NR Peak duration of activated partial thromboplastin time (APTT), mean sec (SD): Heparin: 61.1 (±30.8) No heparin: 66.8±36.8 95% CI difference between means: -11.8-23.3 p = NR Lowest platelet count, x10⁹1⁻¹: Heparin: 172 (±109) No heparin: 156 (±101) 95% CI difference between means: -66.6-35.2 p = NR Phlebitis, n (%): Heparin: 3/35 (8.6) No heparin: 6/31 (19.4) RR: 0.4 (95% CI: 0.1-1.6) p = NR Bleeding, n: Heparin: 2/35 No heparin: 4/31 p = NR Thrombocytopenia, n: |
| 1 | | | | • Heparin: 3/35 |

| Study Information | Population and Setting | Intervention | Definitions | Results |
|-------------------|------------------------|--------------|-------------|---|
| | | | | • No heparin: 4/31 • p = NR |
| | | | | Mortality, n (%): • Heparin: 4/35 (11.4) • No heparin: 6/31 (19.4) • RR: 0.6 (95% CI: 0.2 - 1.9) • p = NR |

Table 66 Risk of Bias for Randomized Controlled Trials on Anticoagulant Infusion

| Author Year | Described as randomized | Randomization appropriately performed | Described as double-blind | Outcome assessor blinded | Study participant blinded | Investigator blinded | Attrition described | Attrition smaller than 10-15% of assigned patients | Attrition appropriately analyzed | Funding source(s) disclosed and no obvious conflict of interest | Overall Risk of Bias |
|------------------------------|-------------------------------|---------------------------------------|---------------------------|--------------------------------|---------------------------------|-------------------------|------------------------|---|--|---|-------------------------|
| Birch 2010 ⁵² | ✓ | ✓ | ✓ | ✓ | ✓ | ✓ | ✓ | √ | ✓ | ✓ | Low |
| Uslu 2010 ⁵⁵ | ✓ | | ✓ | √ | √ | √ | √ | | ✓ | | Moderate |
| Shah 2007 ⁵⁴ | ✓ | ✓ | ✓ | ✓ | ✓ | ✓ | ✓ | √ | ✓ | ✓ | Low |
| Kamala 2002 ⁵³ | ✓ | | ✓ | ✓ | √ | ✓ | ✓ | ✓ | ✓ | ✓ | Low |

D. Evaluation of Study-level Risk of Bias

D.1. Randomized Controlled Trial Checklist

- 1. Described as randomized
- 2. Randomization appropriately performed
- 3. Described as double-blind
- 4. Outcome assessor blinded
- 5. Study participant blinded
- 6. Investigator blinded
- 7. Attrition described
- 8. Attrition smaller than 10-15% of assigned patients
- 9. Attrition appropriately analyzed
- 10. Funding source(s) disclosed and no obvious conflict of interest

D.2. Observational Study Checklist

- 1. Were all study groups derived from similar source/ reference populations?
- 2. Was attrition not significantly different across study groups?
- 3. Was the measure of exposure valid?
- 4. Was the measure of outcome valid?
- 5. Were investigators blinded to endpoint assessment or are the Outcome Definitions objective?
- 6. Were potential confounders identified?
- 7. Were statistical adjustments done for potential confounders?
- 8. Were funding source(s) disclosed and no obvious conflict of interest?

D.3. Descriptive Study Checklist

- 1. Did the study enroll all suitable patients or consecutive suitable patients within a time period?
- 2. Was the study prospectively planned?
- 3. Were independent or blinded assessors used to assess subjective Outcome Definitions?
- 4. Was the study's funding derived from a source that would not benefit financially from results in a particular direction?

D.4. Rating for Overall Risk of Bias

- The risk of Bias was rated as follows:
 - Observational studies:
 - High Risk of Bias: studies with ≤ 50% of checklist items reported
 - Moderate Risk of Bias: studies with > 50% and < 75% of checklist items reported
 - Low Risk of Bias: studies with ≥ 75% of checklist items reported
 - Descriptive Studies
 - High Risk of Bias: studies with ≤ 50% of checklist items reported
 - Moderate Risk of Bias: studies with > 50% of checklist items reported

D.5. Aggregate Risk of Bias

• When the risk of bias was rated as "High" for ≥75% of studies making up the evidence base for a given outcome, one point was deducted for Study Quality in the GRADE table.

E. HICPAC Recommendation Categorization Scheme (2019)

Table 67 Strength of Recommendations

| Strength | Definition | Implied Obligation | Language |
|-------------------------------|---|---|--|
| Recommendation | A Recommendation means that we are confident that the benefits of the recommended approach clearly exceed the harms (or, in the case of a negative recommendation, that the harms clearly exceed the benefits). In general, Recommendations should be supported by highto moderate-quality evidence. In some circumstances, however, Recommendations may be made based on lesser evidence or even expert opinion when high-quality evidence is impossible to obtain, and the anticipated benefits strongly outweigh the harms or when then Recommendation is required by federal law. | A Recommendation implies that healthcare personnel/healthcare facilities "should" implement the recommended approach unless a clear and compelling rationale for an alternative approach is present. | The wording of the Recommendation should specify the setting and population to which the Recommendation applies (eg, adult patients in intensive care unit settings). • Action verbs, eg, use, perform, maintain, replace • Should, should not • Recommend/ is recommended, recommend against/ is not recommended |
| Conditional Recommendation | A Conditional Recommendation means that we have determined that the benefits of the recommended approach are likely to exceed the harms (or, in the case of a negative recommendation, that the harms are likely to exceed the benefits). Conditional Recommendations may be supported by either low-, moderate- or high-quality evidence when: • there is high-quality evidence, but the benefit/harm balance is not clearly tipped in one direction • the evidence is weak enough to cast doubt on whether the recommendation will consistently lead to benefit | A Conditional Recommendation implies that healthcare facilities/ personnel "could," or could "consider" implementing the recommended approach. The degree of appropriateness may vary depending on the benefit vs. harm balance for the specific setting. | The wording of the Conditional Recommendation should specify the setting and population to which the Conditional Recommendation applies when relevant, including: - select settings (eg, during outbreaks) - select environments (eg, ICUs) - select populations (eg, neonates, transplant patients). |

| Strength | Definition | Implied Obligation | Language |
|-------------------|---|--------------------|---------------------------------------|
| | • the likelihood of benefit for a specific patient population or clinical | | Consider |
| | situation is extrapolated from relatively high-quality evidence | | Could |
| | demonstrating impact on other patient populations or in other | | May/ may consider |
| | clinical situations (eg, evidence obtained during outbreaks used to support probable benefit during endemic periods) | | |
| | the impact of the specific intervention is difficult to disentangle | | |
| | from the impact of other simultaneously implemented interventions (eg, studies evaluating "bundled" practices) | | |
| | there appears to be benefit based on available evidence, but the benefit/harm balance may change with further research | | |
| | benefit is most likely if the intervention is used as a supplemental measure in addition to basic practices | | |
| No Recommendation | No Recommendation is made when there is both a lack of pertinent | n/a | "No recommendation can be made |
| | evidence and an unclear balance between benefits and harms. | | regarding" |

Table 68 Justification for Choice of Recommendation Strength

| Components | What to include | Comments |
|-------------------------------------|---|---|
| Supporting Evidence | List the number and type(s) of available evidence used. | eg, " 10 observational studies" |
| Level of Confidence in the Evidence | Level of confidence is low/moderate/high (See Table 3). | eg, "The level of confidence in this evidence is low, as observational studies are at increased risk of bias" |
| Benefits | List the favorable changes in Outcome Definitions that would likely occur if the Recommendation were followed. | Be explicit, clear about pros/cons |
| Risks and Harms | List the adverse events or other unfavorable Outcome Definitions that may occur if the Recommendation were followed. | Be explicit, clear about pros/cons |
| Resource Use | Describe (if applicable) direct costs, opportunity costs, material or human resources requirements, facility needs, etc, that may be associated with following the Recommendation. | HICPAC does not perform its own cost analyses and is not obliged to address cost if analyses are not available and no useful statements can be made. State clearly if information on resource use is lacking. |
| Benefit-Harm Assessment | Classify as "preponderance of benefit over harm" (or vice versa) or a "balance of benefit and harm." Description of this balance can be from the individual patient perspective, the societal perspective, or both. | Recommendations are possible when clear benefit is not offset by important harms or costs (or vice versa); conversely, when the benefit is small or offset by important adverse factors, the balance between benefit and harm prevents a Recommendation. |
| Value Judgments | Summarize value judgments used by the group in creating the Recommendation; if none were involved, state "none." | Translating evidence into action often involves value judgments, which include guiding principles, ethical considerations, or other beliefs and priorities. Stating them clearly helps users understand their influence on interpreting objective evidence. |
| Intentional Vagueness | State reasons for any intentional vagueness in the Recommendation; if none was intended, state "none." | Recommendations should be clear and specific, but if the group chooses to be vague, acknowledging their reasoning clearly promotes transparency. Reasons for vagueness may include insufficient evidence; inability to achieve consensus among panel regarding evidence quality, anticipated benefits/harms, or interpretation of evidence; legal considerations; economic reasons; ethical/religious issues. |

| Components | What to include | Comments |
|------------|--|----------|
| Exceptions | List situations or circumstances in which the Recommendation | |
| | should not be applied. | |

Table 69 Aggregate Level of Confidence in Effect Estimate*

| High | Highly confident that the true effect lies close to that of the estimated size and direction of the effect. For example, confidence in the evidence is rated as "High" when there are multiple studies with no major limitations, there are consistent findings, 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. For example, confidence in the evidence is rated as "Moderate" when there are only a few studies and some have limitations but not major flaws, there is some variation between study results, 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. For example, confidence in the evidence is rated as "Low" when supporting studies have major flaws, there is important variation between study results, the confidence interval of the summary estimate is very wide, or there are no rigorous studies. |

^{*}Based on Grades of Recommendation, Assessment, Development, and Evaluation (GRADE) and the Canadian Task Force on Preventive Health Care

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G. Acronyms and Abbreviations

| Acronym | Expansion |
|-----------|---|
| * | Critical outcome by which decisions are made |
| BSI | Bloodstream Infection |
| CDC | Centers for Disease Control and Prevention |
| CRBSI | Catheter-Related Bloodstream Infection |
| CLABSI | Central Line-Associated Bloodstream Infection |
| CHG | Chlorhexidine Gluconate |
| CoNS | Coagulase-Negative Staphylococci |
| DES | Descriptive Study |
| FDA | Food and Drug Administration |
| GRADE | Grading of Recommendations Assessment, Development and Evaluation |
| HHS | (United States Department of) Health and Human Services |
| HICPAC | Healthcare Infection Control Practices Advisory Committee |
| IV | Intravenous |
| MRSA | Methicillin-Resistant Staphylococcus aureus |
| MSSA | Methicillin-Sensitive Staphylococcus aureus |
| NICU | Neonatal Intensive Care Unit |
| OBS | Observational Study |
| PICC | Peripherally Inserted Central Catheter |
| PCR | Polymerase Chain Reaction |
| PI | Povidone Iodine |
| QI | Quality Improvement |
| RCT | Randomized Controlled Trial |
| S. aureus | Staphylococcus aureus |
| TAP | Targeted Assessment for Prevention |
| UAC | Umbilical Arterial Catheter |

| Acronym | Expansion |
|---------|---------------------------|
| UVC | Umbilical Venous Catheter |
| VLBW | Very Low Birthweight |