Microbial colonization research in humans expanded significantly over the last 3 years with the recognition that the composition and differentiation of the human microbiome is related to the health of individuals. The human microbiome influences the host immune system and plays a significant role in acquiring as well as preventing disease states. Humans and microbes form a relationship that starts before birth and evolves throughout the lifespan. High-risk term and preterm infants are some of the most vulnerable to an altered microbiome due to the atypical neonatal intensive care unit (NICU) environment of care following birth.

The human microbiome is defined as the entire collection of naturally occurring bacteria, fungi, and viruses, including their DNA that exists in the human body. See Table 1 for a list of common terms used in the study of the microbiome. The healthy infant’s microbiome begins prenatally and evolves rapidly after delivery as the infant is exposed to the mother’s microbiota. Contact with these initial microbes is beneficial to the health of infants by forming the basis of nutrient utilization, gut barrier function, and immune development. Alterations in the formation of the infant microbiome can occur and are associated with adverse gut barrier function and immune response. Infants born via cesarean section have microbial flora similar to environmental microbes, whereas infants born vaginally have intestinal microbial content similar to the atypical neonatal intensive care unit (NICU) environment of care following birth.

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the mother’s vaginal and intestinal flora including the commensal bacteria Bacteroides, Bifidobacterium, and Escherichia coli. 12,13 This difference in microbial composition is important because commensal bacteria facilitate the development of the infant’s immune system. 9 Infants born by caesarean section may take up to 6 months of age to colonize the commensal bacteria. 2 Epidemiological data show that infants born via caesarean section are also more prone to future childhood disease such as allergic rhinitis, asthma, and celiac disease. 8 Following delivery infants continue to develop and alter their microbiome on the basis of the type of feeding, contact with microbes from the environment, and from economical changes such as improved sanitation, increased antibiotic usage, and immunization prophylaxis. 14,15

Although the mode of delivery provides the foundation for the infant’s initial microbiota, development of the microbiome continues with ongoing interaction with the mother, family, and home environment. Unfortunately, high-risk infants are separated from their mothers following delivery and for weeks to months are cared for in the complex environment of the NICU. Every NICU has its own genera of microbes, which can include commensal and pathogenic organisms. 16,17 With the NICU genera as the context for microbiome development, the infant’s ongoing interaction with multiple healthcare providers and caregiving processes has the potential to impact microbial colonization. A recent study showed that the risk of acquiring nasal colonization with coagulase-negative Staphylococcus for all neonates admitted in a NICU was 55.9%. 18 High-risk term and preterm infants are at an increased risk of infection due to their illness, and in preterm infants due to immature skin and immune systems. 19,20 Caregiving equipment not only comes in contact with infants, but items like electrocardiogram (EKG) leads, monitoring probes, adhesive dressings, and

| TABLE 1. Common Terms in the Microbiome Literature |
| --- | --- | --- |
| **Keyword** | **Definition** | **Examples** |
| 1. Human microbiome | The entire collection of naturally occurring bacteria, fungi, and viruses, including their DNA, that exists in the human body, which are critical for normal development and health. | Bacteria locations: mouth (anaerobes); large intestine (aerobes and anaerobes); skin (predominately aerobes) | Virus locations: implant into living cells mainly in the respiratory (coxackie viruses), gastrointestinal (adenovirus), skin-penetrating, and genital routes (human papillomavirus) | Fungi locations: superficial including the skin, hair, and nails (candidiasis) |
| 2. Microbiota | Complex communities composed of many microorganisms including bacteria, fungi, and viruses that colonize various sites in the human body. | | | |
| 3. Genera | The usual subdivision of a family or subfamily in the taxonomic classification of organisms, usually consisting of more than 1 species. | Eq, Enterobacter (genus Enterobacter) is a group of rod-shaped bacteria of the family Enterobacteriaceae, gram-negative facultative anaerobes | Eq, Escherichia coli (genus Escherichia) is a gram-negative, facultative anaerobic, rod-shaped bacterium |
| 4. Microbes | Microorganisms such as bacteria, fungi, and viruses that can perform beneficial and harmful effects. | | | |
| 5. Commensal microbes | Organisms that produce positive effects for a host. Indigenous: present on body surfaces covered by epithelial cells and exposed to the external environment. | Bacteroides, Bifidobacteria, Enterobacteriaceae, Enterococcaceae Escherichia coli, Lactobacilli Neisseria (all except N. gonorrhea and N. meningitides, Staphylococci spp., Streptococci spp.) |
| 6. Pathogenic microbes | Organisms that produce negative effects for a host. Indigenous organisms may induce disease when the host is compromised or with commensal microbe overgrowth. | Acinetobacter, Actinobacteria, Citrobacter Clostridium spp., Firmicutes, Gemella, Geobacillus, Halomonas, Klebsiella spp. Proteobacteria, Pseudomonas aeruginosa, Shewanella, Ureaplasma (commensal in sexually active women) | |
chemical burns that can damage the epithelium can also lead to increased risk of microbial invasion.20 Over the past 2 decades several NICU outbreaks of *Pseudomonas aeruginosa* and *Klebsiella pneumoniae* neonatal infections were linked to healthcare providers and caregiving activities,21-24 but they fail to inform us about the development of the infant’s microbiome. Understanding the colonization patterns of microbes on the skin and in the gastrointestinal tract of infants can help NICU caregivers develop interventions to promote infant health. This systematic review of the literature summarizes what is known about the impact of the NICU environment on microbial colonization of high-risk term and preterm infants and identifies implications for clinical practice and future research.

**THE MICROBIOME**

Microbiota are complex communities composed of many microorganisms including bacteria, fungi, and viruses that colonize various sites in the human body.1 In fact, most of the body’s cells are not human but are microbial in origin with “an estimated microbial/human ratio of 10:1.”2 Of all sites, the human gut houses 10 times as many microorganisms as other areas of the body.7 The skin, bladder, mouth, and vagina are also heavily colonized sites.25

All microbes, which are microorganisms such as bacteria, viruses, and fungi,1 were once thought to be harmful. However, microbes perform various positive functions for the host such as regulating the development of the gut, preventing harmful bacteria from colonizing, and synthesizing vitamins and unused substrates.8 Although bacteria can produce positive effects for a host, a fine line exists between acquiring commensal bacteria and acquiring pathogenic bacteria.23 Pathogenic bacteria can be introduced through a multitude of factors such as altered host immune function, genotype, diet, and the environment.25

The Human Microbiome Project, sponsored by the National Institute of Health, was developed to understand what determines a healthy versus an unhealthy aggregate of genetic material of bacteria, fungi, and viruses and to identify microbial flora that exist in the different areas of the body.10 Current research aims to understand how the microbiome forms and what factors, such as the environment, contribute to commensal and pathogenic microbial colonization. How the microbiome relates to neonatal health is of importance due to high-risk and preterm infants’ increased risk of infection in the environment of the NICU, as well as their risk for altered health and developmental outcomes. The acquisition of the microbiome during infancy can influence immune health over the lifespan.26

This systematic review describes the relationship between factors within the NICU environment and microbial colonization of high-risk infants. For the purpose of this review, environment was defined as the physical environment of care and caregiving practices. The findings will increase healthcare professionals’ awareness of the potential impact of the NICU environment on skin and gut colonization of infants and identify caregiving interventions to improve infant outcomes.

**LITERATURE SEARCH METHODS**

**Data Sources**

Prefered Reporting Items for Systematic Reviews and Meta-Analyses standards were used as a framework for this review.27 Electronic databases used to identify relevant articles in English with an available full-text or print copy included PubMed BIOSIS Citation Index, Google Scholar, the Cumulative Index for Nursing and Allied Health Literature, and BioMedSearcher. Search terms used to identify research articles relevant to the research question included *microbiome*, *microbiota*, *microbe*, *antibiotic*, *infant skin care*, *isolate*, *delivery mode*, *environment*, *infection control*, *immunity*, *incubator*, *necrotizing enterocolitis* (NEC), *infant*, *premature emollients*, *nursery*, *NICU*, and *gut colonization*. No limits were set on research articles regarding the date of publication or country of origin.

**Study Selection**

A literature search was conducted in January 2014, updated in June and November 2014, and revised in January 2015. Figure 1 summarizes the study selection. After screening titles and abstracts of 250 articles, 39 articles were chosen to read thoroughly on the basis of discussion of the environmental influence of microbial colonization on infants. Eleven articles were chosen as sources for this review on the basis of our definition of NICU environment and potential environmental influences on the microbiome of high-risk and preterm infants. These articles included literature about the influence of the mother and mode of delivery on the infant, factors that influence infant gut colonization, and the risks associated with the colonization of the NICU environment.

**Data Extraction**

Each article was read carefully and categories were identified that had the potential to influence the infant microbiome. The literature analysis presented challenges given the emerging science about the infant microbiome and the identification of few studies specific to the relationship between the physical NICU environment or caregiving practices on the
FIGURE 1

Records identified through database searching (n = 290)

Additional records identified through other sources (n = 10)

Records after duplicates removed (n = 250)

Records screened (n = 250)

Records excluded (n = 190)

Full-text articles assessed discussed environmental influence of microbial colonization (n = 39)

Full-text articles excluded included single case studies of healthy infants, no relationship to NICU environment, opinion and review articles (n = 28)

Studies included in the synthesis (n = 11)

Search strategy and summary.

infant microbiome. As evidence of the developing nature of this work, the 11 articles chosen as sources for this review were all published within the last 4 years (2011-2014). The study designs included 6 cohort studies, 2 longitudinal studies, 1 retrospective descriptive study, 1 prospective surveillance study, and 1 descriptive comparative study. No randomized controlled trials or systematic reviews were found related to the potential environmental influences on the infant microbiome.

Definitions of the microbiome, microbiota, and microbe were used interchangeably in many studies. Consistent use of these terms would make research on this body of knowledge stronger and would decrease variability on this subject.

DATA SYNTHESIS

Categories of factors associated with changes in the infant’s microbiome in the 11 articles were parental skin, feeding (human milk vs formula), environmental surfaces, nursing workspaces, and infant caregiving equipment (e.g., ventilators, incubators, and warmers); healthcare provider skin; and antibiotic use (see Table 2).

Parental Skin

Many NICUs encourage skin-to-skin contact, consisting of “chest-to-chest placement of the infant with the mother at an incline of 30-40 degrees.” Skin-to-skin care (SSC) allows for positive touch between the infant and mother as well as transfer of microbes. No studies were found that examined the relationship between skin-to-skin contact and the infant microbiome; however, 1 study did examine infection rates after mother skin-to-skin care in preterm infants. After 5 consecutive days of skin-to-skin care for 90 minutes, only 1 infant developed a hospital-acquired infection within 7 days after the skin-to-skin care ended. None of the mothers had any signs or symptoms of infection during the first 30 days postdischarge.

Two studies identified microbial transfer from parents to infants cared for in the NICU. In a NICU with an outbreak of Staphylococcus aureus (S. aureus), S. aureus isolates were identified from 2 fathers and the nipples of 1 mother prior to any symptoms of infection in the infant. A second study showed that organisms found in the NICU environment were also found on parents hands and in the stools of 2 infants.
<table>
<thead>
<tr>
<th>Title/Authors (Y)/Country</th>
<th>Population/Study Design</th>
<th>Environmental Factors</th>
</tr>
</thead>
<tbody>
<tr>
<td>Effect of skin-to-skin contact on preterm infant skin barrier function and hospital-acquired infection Abouelfettoh et al (2011)28 United States</td>
<td>Pretest-test-posttest cohort study N = 10 preterm infants with 90 min of SSC daily for 5 d Blood cultures results during hospitalization and up to 4 wk postdischarge Signs and symptoms of infection after discharge were defined as positive maternal response to 1 or more of a 3-question questionnaire concerning infection after discharge from the NICU Parental skin One infant was blood culture positive 7 d after SSC No infants had signs of infection within 4 wk of discharge</td>
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<tr>
<td>Microbes in the neonatal intensive care unit resemble those found in the gut of premature infants Brooks et al (2014)29 United States</td>
<td>Longitudinal descriptive 2 VLBW (&lt;1500 g) infants cared for in the same area for first month of life Stool samples collected every 3 d 33 swabs were collected each room surfaces (sink, feeding and intubation tubing, hands of healthcare providers and parents, general surfaces, nurse station computer keyboard and mouse, and cell phone Environmental surfaces, caregiving equipment, healthcare provider skin, parental skin Distinctly different GI tract colonizations in the 2 infants Gut organisms were widely distributed throughout the room environment— included <em>Staphylococcus epidermidis, Klebsiella pneumoniae, Bacteroids fragilis, and Escherichia coli</em> <em>Klebsiella pneumoniae</em> in infant 1 and <em>Finegoldia magna</em> in infant 2 were present in the room prior to detection in the gut Keyboards, mouse, and telephones had the lowest amount of colonizing organisms detected in the gut Intubation and feeding tubing had the highest amount of colonizing organisms detected in the gut</td>
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<tr>
<td>Staphylococcus aureus reservoirs and transmission routes in a Portuguese neonatal intensive care unit: A 30-mo surveillance study Conceicao et al (2012)30 Portugal</td>
<td>Prospective surveillance 16 clinically symptomatic infants who were <em>Staphylococcus aureus</em> hemo positive; mean gestation age of 31 wk Swabs from the infants’ parents nares; NICU healthcare workers nares during the 4 d preceding each infection case; mothers’ nipples in case of breast feeding; NICU environment (cardiorespiratory monitors, incubators, milk pumps, stethoscopes, plastic folder protecting the infants clinical records hanging on the incubators, and telephones) Environmental surfaces and caregiving equipment, parental skin, health care provider skin Mean time to acquire an <em>S. aureus</em> infection after admission was 15 d (ranging from 2 to 39 d) <em>S. aureus</em> isolates recovered from infant’s hemocultures, catheters, umbilical exudates, wounds, pus of intestinal abscesses, pus of skin abscesses, and urine MSSA isolates recovered from plastic folder in the NICU environment Infection cases had the same strain recovered from the infant and other sources such as at least one health care worker, the environment (plastic folder), mother’s nipples</td>
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<tr>
<td>Bacterial community structure and functional contributions to emergence of health or necrotizing enterocolitis in preterm infants Claud et al (2013)31 United States</td>
<td>Descriptive cohort study 10 Preterm infants Stool samples of 5 infants with NEC and 5 healthy matched controls Feeding type Control fecal samples had a temporal pattern of microbiota typical of healthy full-term breast-fed infants, Microbiota development in NEC patients diverged from healthy controls beginning three weeks prior to diagnosis</td>
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### TABLE 2. Potential Environmental Influences on the Infant Microbiome, Continued

<table>
<thead>
<tr>
<th>Title/Authors (Y)/Country</th>
<th>Population/Study Design</th>
<th>Environmental Factors</th>
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<tbody>
<tr>
<td><strong>Intestinal microbial ecology and environmental factors affecting necrotizing enterocolitis</strong>&lt;br&gt;Torrazza et al (2013)&lt;sup&gt;32&lt;/sup&gt;&lt;br&gt;United States</td>
<td>Descriptive case controlled cohort&lt;br&gt;N = 53 preterm infants ≤32 wk GA; 18 NEC cases; 35 control cases&lt;br&gt;3 NICUs (1 in Gainesville, Florida, 2 in Jacksonville, Florida)&lt;br&gt;Stool samples at 2, 1, and 0 wk, prior to the diagnosis of NEC. Samples from matched control infants were chosen during the same week of life at which the samples from the cases were obtained</td>
<td>Environmental surfaces, feeding type, antibiotic use&lt;br&gt;Microbiota composition differed between the 3 NICUs where antibiotic use also differed. Incidence of NEC was 12.4% for hospital 1 vs 6.8% for hospital 2 and 3&lt;br&gt;Control infants received human milk 57.1% of the time while infants with NEC received human milk only 27.8% of the time&lt;br&gt;Abnormal patterns of colonization with predominance of Proteobacteria early in life or later in the days closer to the development of NEC may be associated with a greater risk of developing NEC</td>
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<td><strong>Development of the preterm gut microbiome in twins at risk of developing necrotizing enterocolitis and sepsis</strong>&lt;br&gt;Stewart et al (2013)&lt;sup&gt;33&lt;/sup&gt;&lt;br&gt;Spain</td>
<td>Longitudinal study&lt;br&gt;27 infants (12 twin pairs and 1 triplet set)&lt;br&gt;173 stool samples&lt;br&gt;18 expressed breast milk samples from 4 mothers</td>
<td>Feeding type, antibiotics&lt;br&gt;Staphylococcus, Corynebacterium, and Propionibacterium were found in high abundance in the gut microbiome&lt;br&gt;Gut microbiome was more similar between siblings than unrelated individuals&lt;br&gt;Reduction in diversity and increasing dominance of Escherichia sp. preceded NEC was not observed in the healthy twin&lt;br&gt;Antibiotic treatment reduced Escherichia sp. and increased other Enterobacteriaceae in the gut</td>
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<td><strong>Surface microbes in the neonatal intensive care unit: Changes with routine cleaning and over time.</strong>&lt;br&gt;Bokulich et al (2013)&lt;sup&gt;16&lt;/sup&gt;&lt;br&gt;United States</td>
<td>Pre-post cohort study&lt;br&gt;Environmental cleaning NICU surface samples; N = 147&lt;br&gt;Neonate-associated (eg, incubator, pacifier, and ventilator)&lt;br&gt;Room environment (eg, computer, space bar, and cardiac monitor)</td>
<td>Environmental surfaces&lt;br&gt;Fungal populations included Saccharomycyces cerevisiae, Cryptococcus albidus, Debaryomyces fabryi, and Candida albicans and were less prominent on neonatal than room surfaces&lt;br&gt;Cleaning with antibacterial wipes significantly reduced the total microbial load on surfaces and the collection of microbiota shifted to primarily nonpathogenic organisms&lt;br&gt;Levels of common NICU enteric genera were not altered by cleaning (Enterococcus, Klebsiella, Escherichia, and Pseudomonas)</td>
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<tr>
<td><strong>Cold spots in neonatal incubators are hot spots for microbial contamination</strong>&lt;br&gt;de Goffau et al (2011)&lt;sup&gt;34&lt;/sup&gt;&lt;br&gt;Netherlands</td>
<td>Prospective cohort study&lt;br&gt;23 Calo incubators&lt;br&gt;Group 1 N = 39 hot; 50 cold spots samples&lt;br&gt;Group 2 N = 40 hot; 60 cold spots samples&lt;br&gt;At infant transfer to a clean incubator after 4-7d of use&lt;br&gt;Swab samples from 4 hot and 5 cold spots</td>
<td>Environmental surfaces and caregiving equipment&lt;br&gt;Higher average incubator temperatures (≥34°C) and relative humidity values (≥60%) were associated with higher levels of microbial contamination at cold spots vs hot spots&lt;br&gt;Primary organism found was Staphylococci</td>
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(continues)
Feeding Type

The impact of breast milk and formula on the developing microbiome of premature infants has been hypothesized especially in the development of NEC. The relationship between feeding type was addressed in 3 studies that explored the differences in the gut microbiome of premature infants who developed NEC versus the gut microbiome of healthy control premature infants who did not develop NEC. In 1 study, stool samples of NEC cases showed a high proportion of Proteobacteria (61%) at 2 weeks and Actinobacteria (3%) at 1 week before diagnosis with NEC, along with lower amounts of Bifidobacteria and Bacteroidetes. Stool samples of control infants who did not acquire NEC had less abundance of Proteobacteria (19%) and Actinobacteria (0.4%), and they received human milk 57.1% of the time compared with only 27.8% of the time for infants who developed NEC. A second group of infants had stool samples dominated by Proteobacteria with members of the Firmicutes; however, prior to 2 weeks old, infants who later developed NEC had fewer numbers of Firmicutes compared with infants who did not develop NEC. The control infants’ stool samples also had a microbiota similar to healthy full-term, breast-fed infants. Finally, in a related study, 173 stool samples were obtained from 12 sets of twins and 1 set of triplets, as well as 18 samples of expressed breast milk from 4 mothers. Both the expressed breast milk and stool samples shared a common microbiome composed of Enterobacteriaceae, Enterococcaceae, and Staphylococcaceae, although both had a low abundance of Bifidobacteria and Lactobacilli. The expressed breast milk samples had an evolving microbiome throughout lactation as new diversified bacteria continually arose.

Environmental Surfaces, Nursing Workspaces, and Caregiving Equipment

Preterm infants are immunocompromised, making them highly susceptible to hospital-acquired infections. Virtually every object and piece of equipment used in the NICU environment can serve as a reservoir for microbial contamination. Four studies examined several aspects of the NICU environment.
in an effort to better understand how they might impact the infant microbiome.16,29,30,34 Specific measures utilized within these studies included sampling before and after cleaning with antibiotic wipes,16 cold versus warm spot sampling in incubators,14 predominant NICU organisms living within specific nurseries regardless of routine cleaning and hand washing,29 and lack of hand washing and routine cleaning before and after implementing standard precautions.30 Microbiome samples of environmental surfaces revealed complex and diverse environments and included species associated with nosocomial infections common in the neonatal population.16,29,34 Surfaces that came in direct contact with infants (continuous positive airway pressure machines, stethoscopes, ventilators, incubators, and radiant warmers) were colonized with Streptococcus, Staphylococcus, Neisseria, and Enterobacteriaceae.16,30 whereas room environment surfaces (computer screens, a computer mouse, freezer handles, hand sanitizer bottles, door handles, telephones, heart rate monitor screens, and alarm buttons) were colonized with communities of Geobacillus, Halomonas, Shewanella, Acinetobacter, and Gemella.16 Over time the surfaces with the greatest variation were those exposed to human skin.29

Like other surfaces that come into direct contact with the infant, the microbiome of incubators included Streptococcus,16 Staphylococcus,16,30 Neisseria, and Enterobacteriaceae.16 The uniformity of heat and humidity was related to the colonization of incubators. In humidified Caleo (Dräger, Drägerwerk AG & Co. KGaA, Telford, PA) incubators with average temperatures 34°C or more and relative humidity levels 60% or more, microbial contamination with Staphylococcus was significantly higher at incubator cold spots than at hot spots.14 The cold spots were found at the front, back sides, and middle of the incubator where the humidified air did not reach as well. Hot spots were closer to the hot air vents. Across a wide variety of surfaces including incubators, ventilators, monitors, and computers, cleaning with disinfectant wipes significantly decreased the bacterial and fungal load on surfaces sampled before and after cleaning.16

Specific infant caregiving supplies like other NICU surfaces had diverse microbiome communities. Pacifiers were colonized with diverse communities of Streptococcus, Staphylococcus, Neisseria, and Enterobacteriaceae,16 and stool samples of 2 infants were similar to gut organisms (Staphylococcus epidermidis, Klebsiella pneumonia, Bacteroids fragilis, and E. coli) prevalent in intubation and feeding tubing.29

The relationship between the general NICU environment and the infant gut microbiome was explored in 4 studies.22,25,37 Stool colonization patterns in neonates revealed that the diversity of clostridia species increased throughout hospital stay, which included Clostridium perfringens, Clostridium butyricum, Clostridium difficile, and Clostridium paraputrificum.35 Different fecal microbiota composition also existed in neonates housed in different NICUs; 1 hospital had more colonization with Bacteroidetes and Proteobacteria versus colonization with Firmicutes at another hospital.32 Stool samples of infants who acquired NEC showed a higher portion of Proteobacteria and Actinobacteria and lower numbers of Bifidobacteria and Bacteroidetes.32 Gastrointestinal aspirates of very low birth-weight infants revealed that Staphylococci were dominant overall, Ureaplasma were dominant in the first week of life, and by the fourth week of life, gram-negative bacteria increased.34 Firmicutes were present in the majority of the neonates and only 2 species of S. epidermidis were found.16 When stool, skin, and saliva samples were compared among low birth-weight infants, Klebsiella/Enterobacter, Enterococcus, and Citrobacter were present in stool samples; Staphylococcus was present on the skin, and Streptococcus was present in the saliva.17 However, when stool samples of low birth-weight infants were compared with stool samples of normal birth-weight infants, Escherichia was highly present in the stool of normal birth-weight infants but absent in low birth-weight infants who had an abundance of Enterobacter, Enterobacteriaceae, Enterococcus, and Staphylococcus.17 These findings were related to delay in enteral feedings and increased length of hospital stay in preterm infants.37 Finally, when samples were compared with adult microbiota, preterm skin microbiota was most like adult skin microbiota; preterm saliva and stool microbiota were the least like adult microbiota.

Healthcare Provider Skin
The mechanisms of human-to-human transfer of organisms are well known and are 1 source of nosocomial infections. Two studies looked at direct transfer of microbes from healthcare providers’ skin to infants cared for in the NICU.29,30 From 154 surveillance screening swabs in a NICU with an outbreak of S. aureus, 24 S. aureus isolates were identified from 18 healthcare workers.31 This bacteria was identified prior to the symptoms occurring in the infants cared for by the providers with S. aureus isolates.30 With more than 50% (n = 64) healthcare providers screened, a 28% prevalence of S. aureus colonization was identified.30 In a second study, samples from healthcare providers skin revealed Actinobacteria, Firmicutes, and Proteobacteria.29 Caregiver hands were also contributors to the variation of environmental organisms present on environmental surfaces and caregiving supplies. Over time the stools of 2 infants cared for in the same nursery developed similar microbes.29
Antibiotic Use

Antibiotic use in NICUs remains widely diverse and controversial. Antibiotics are effective in depleting pathogenic bacteria during disease states but simultaneously deplete commensal bacteria. Depleting commensal bacteria negatively alters the infant microbiome and is thought to increase the risk of acquiring NEC. The relationship between antibiotic use, gut microbes, and NEC was explored in 3 articles. One study specifically tested for Clostridia strains, which were susceptible to various antibiotics including amoxicillin-clavulanic acid, piperacillin-tazobactam, chloramphenicol, metronidazole, linezolid, and vancomycin. Strains were resistant to clindamycin, cefotaxime, tetracycline, and moxifloxacin. Antenatal antibiotic treatment was associated with significantly decreased Clostridia levels, whereas intrapartum antibiotic therapy was unrelated. Neonatal antibiotic therapy was also associated with decreased levels of Clostridial colonization when given for more than 10 days. In a second study, prior to the development of NEC, infants were exposed to different antibiotics and varying durations of treatment at 3 different NICUs. Antibiotic usage was associated with observed differences in the infants’ microbiome. Infants who went on to develop NEC had a higher proportion of Proteobacteria (61%) and Actinobacteria (3%) at 2 weeks and 1 week prior to the diagnosis, respectively. The infants who developed NEC also had lower numbers of Bifidobacteria counts and Bacteroidetes. The final study, antibiotic therapy was given to a set of twins and showed reduced Escherichia sp. and increased amount of Enterobacteriaceae in stool samples, which reversed in 1 twin when antibiotics were stopped. The other twin developed NEC and received subsequent antibiotic therapy, which altered the bacterial community from the sibling, showing increased levels of Klebsiella sp. and smaller increases in members of the Bifidobacteriaceae family.

DISCUSSION

The limited studies in this review were not able to answer the complex sequencing questions necessary to determine exactly when and from what the infant acquires its microbiome. However, specific species from the NICU environment are associated with nosocomial infections, and infant complications like NEC reveal patterns of gut microbiota development different from infants who do not develop NEC. Researchers continue to define what constitutes the normal microbiome for healthy term infants, and research techniques in this area are advancing rapidly. What is known about the microbiome in preterm infants remains obscure, and much research is still warranted. The recent literature provides evidence about the diversity of the microbiome in the NICU environment. However, exactly how the microbiome of the NICU environment, caregivers, feeding type, and antibiotics affect the development of the infant’s microbiome over time requires further exploration.

Parental contributions to the healthy infant’s microbiome are typically viewed as positive such as in the practice of skin-to-skin care and breastfeeding. However, the typical parental microbiome may also be altered as they spend significant amounts of time in the NICU environment. Several studies identified bacterial and fungal colonization of NICU surfaces, equipment, skin and caregiving supplies that are associated with nosocomial infections in the neonatal population. Intensive care nurseries commonly engage in infection control bundles and routine cleaning of bedside and nursery equipment as a method to prevent nosocomial infections, and at least 1 study demonstrated a decrease in bacterial communities following routine cleaning practices. However, how the routine cleaning might affect colonization of both commensal and pathogenic bacteria in infants is unclear.

Humidified incubators, a mainstay piece of equipment in the treatment of preterm infants in NICUs, are beneficial for skin integrity and temperature regulation, yet the warm moist environment is also a habitat for microbial growth. Staphylococci, a well-known infectious agent in preterm infants and gram-negative bacteria, a risk factor for infection related to neonatal death, were found in the cold areas of humidified incubators. Yet, the frequency with which incubators should be cleaned to minimize risk of infection while maintaining infant safety is unknown. The transfer of infants into a clean incubator is not without associated risks of dislodgement of catheters and equipment, as well as over-stimulation of the infant.

Several studies revealed that the infant gut is colonized differently on the basis of the composition of the NICU environment. Infants appear to acquire organisms in their gut days to weeks after they appear on the surfaces and equipment of the NICU environment. Microbial invasion of the gut from the environment also plays a role in neonates acquiring NEC, although specific pathogens have not been identified. However, infants who developed NEC had markedly different microbiota, which could be attributed to the NICU environment, type of feeding, and antibiotic use. Along with the microbiome, host physiology, such as how an infant metabolizes milk, may also play a role in the development of NEC. Infants who received breast milk had gut microbiota more similar to healthy breastfed term infants, revealing that they may be at a lower risk for developing NEC. Together the findings do not provide definitive evidence to which type of
feeding is most beneficial for the development of the gut microbiome; however, given the many other benefits of breast milk it would appear prudent to use breast milk when available. Finally, while the length and criteria for antibiotic therapy remains controversial, antibiotic use appears to be associated with a change in the pattern of the gut microbiome consistent with the development of NEC.32,33,35

The results of this review did not provide information about the development of the infant’s skin microbiome. Although not studied to date, it is known that the skin provides a protective barrier from microbial invasion.38,39 Infants in the NICU are faced with infection risk from EKG leads, monitoring probes, adhesive dressings, and chemical burns that can damage the epithelium and lead to increased risk of microbial invasion.20 Therefore, skin integrity and knowledge about the skin microbiome in premature and high-risk infants are essential during the first 2 weeks of life when the epidermis is beginning to mature and neonatal infections are common.19,20,38,39

Implications for Practice
Although the findings of this review are unclear about how the NICU environment specifically influences microbial colonization in neonates, the presence of pathogenic bacteria indicates the need for continued diligence in the bundle of practices implemented to reduce nosocomial infections.40,41 Simple techniques such as vigilant hand hygiene, wearing gloves, and practicing sterile technique during invasive procedures can prevent infections especially central-line bloodstream infections60 and should include not only healthcare providers but also parents.

Routine surface cleaning with disinfectant solutions should be employed to decrease colonization of environmental bacteria on surfaces16 as well as incubators.34 Minimizing condensation on the inside of incubators, wiping the vapor away with a clean cloth, and not leaving the cloth inside the incubator for consecutive use may also prevent excessive colonization.34 Finally, it may be helpful to change incubators at least once a week.34

Evidence-based skin care guidelines should also be implemented to protect the epidermal barrier of all infants, but especially preterm infants.42 Standardization of skin care to facilitate microbiome development is lacking, but in at least 1 study cleaning the skin of preterm infants between 28 and 36 weeks of age with 0.25% chlorhexidine within 3 hours of birth reduced axillary skin colonization at 24 hours.19 Although this study did not report any adverse impact on the skin or temperature of the infant, the efficacy of bathing with chlorhexidine is unclear. The same study found that cleaning with saline offered the same reduction in axillary skin colonization.19 The use of topical emollients such as Aquaphor (Beiersdorf Inc, Connecticut) is controversial. Topical emollients have proven to be effective at decreasing transepidermal water loss and maintaining skin integrity in premature infants, but some studies have shown an increased risk of nosocomial infection with emollient use.43 The use of No-Sting (3M, Minnesota) is just as effective as Aquaphor at decreasing transepidermal water loss and maintaining skin integrity, but further studies need to be conducted on its use.44

Labor and delivery nurses as well as NICU nurses should also be strong advocates for new mothers to breastfeed or pump breast milk for their infants. The maternal milk microbiome contains beneficial bacteria for infants to help develop a mature immune system and fight infections.45 Formula preparation does not try to mimic breast milk properties, although it does not contain the beneficial, natural immunoglobulins, oligosaccharides, and fatty acids contained in maternal breast milk.46,47

Practitioners and providers should be cautious in prescribing antibiotics to neonates if the need for antibiotics is not fully justified. Depending on the institution, many high-risk infants will receive antibiotics on admission immediately after birth. Antibiotics will not only deplete pathogenic bacteria but also can be detrimental by eliminating all commensal bacteria.48 Antibiotics deplete the diversity of microbes within the microbiome as well as halt the development of commensal bacteria.49 Antibiotic use has also shown to be associated with the establishment of NEC in premature infants.32

Implications for Future Research
The microbiome of neonates is complex and what is considered to be a normal microbiome is not clear. Additional research should attempt to identify normal microbiome development for not only the gut, but also for the skin of high-risk and preterm infants. Although various infection control bundles have been put in place, the impact on the microbiome is not known. Additional research should evaluate how various caregiving activities such as oral suctioning and the feeding tube type and method may impact the microbiome of the infant’s gut. How long feeding tubes should remain in place and whether in and out placement of feeding tubes promotes or hinders the gut microbiome is unknown. In addition, while breast milk is presumed to be the best for infants, the impact of mother’s own milk versus banked breast milk on the microbiome should be evaluated. Finally, further research is required on the use of probiotics and prebiotics to help form the gut microbiota in infants, as well as determine whether they can help prevent infections and the risk of NEC.

Although current research exists focused on neonatal skin care guidelines, more research should focus on evaluating how disinfectants, bathing, wound care products, and skin protection products like No-Sting and Aquaphor affect the development
of the skin microbiome. Early skin care practices have the potential to not only impact the skin microbiome in the NICU but may also influence the development of future autoimmune problems such as asthma and atopic dermatitis.

Future studies of the impact of the NICU environment on the developing microbiome should include all infants, not just the infants who develop infections. Larger sample sizes should also be utilized in new studies, as all studies used within this systematic review were of small sample sizes. Microbiome samples from the NICU environment, healthcare providers, parents, and infants collected simultaneously over time would significantly add to our understanding of how the microbiome of the infant cared for in the NICU develops. In addition, researchers should use current genomic DNA standardized procedures rather than cultures to evaluate the impact of the NICU environment on the infant. Finally, new research should explore how the infant’s developing microbiome in the NICU impacts the development of the immune system and long-term health outcomes.

**CONCLUSION**

The microbiome of high-risk and preterm neonates may be altered by the environment in various ways beginning with the mode of delivery. Potential influencing factors from the NICU environment include the genera of the NICU itself and essential caregiving contact between the infant and the parents and healthcare providers. Not all of these environmental influences are harmful, yet some were related to an increased risk of obtaining pathogenic microbes. How these factors influence long-term similarities or differences in the microbiome of the infant hospitalized in the NICU compared with infants who are discharged home with their family following delivery remains unknown.

### References


26. Mueller NT, Bakacs E, Combellick J, Grigoryan Z, Dominguez-Bello MB. Advances in Neonatal Care. For more details on manuscript submissions. Please see the author guidelines for young, sick, or genetically impaired are reported. Reports on special programs to provide hospital-acquired infection.


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