Factors associated with progression to infection in methicillin-resistant *Staphylococcus aureus*-colonized, critically ill neonates

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Abstract

Objective To identify factors associated with development of symptomatic infection in infants colonized with methicillin-resistant *Staphylococcus aureus* (MRSA) in the Neonatal Intensive Care Unit (NICU).

Study design This case-control study was performed at St. Louis Children’s Hospital NICU from 2009 to 2019. The MRSA-colonized infants who developed symptomatic MRSA infection (cases) were matched 1:3 with MRSA-colonized infants who did not develop infection (controls). Demographics and characteristics of NICU course were compared between groups. Longitudinal information from subsequent hospitalizations was also obtained.

Results Forty-two infected cases were compared with 126 colonized-only controls. Cases became colonized earlier in their NICU stay, were less likely to have received mupirocin for decolonization, and had a longer course of mechanical ventilation than controls. Longitudinally, cases had a more protracted NICU course and were more likely to require hospital readmission.

Conclusion Progression from MRSA colonization to symptomatic infection is associated with increased morbidity and may be mitigated through decolonization.

Introduction

In hospitalized neonates, *Staphylococcus aureus* is a common cause of late-onset infection, leading to significant morbidity, mortality, and prolonged hospitalization in those who are infected [1]. Methicillin-resistant *S. aureus* (MRSA) infections in neonates include skin and soft tissue infections, respiratory tract infections, pneumonia, bacteremia, osteomyelitis, and septic arthritis [2]. Morbidities that occur as a result of late-onset infection include a prolonged need for mechanical ventilation and subsequent bronchopulmonary dysplasia, necessitating longer hospital courses and decreasing likelihood of survival [1]. A study in a Neonatal Intensive Care Unit (NICU) in North Carolina estimated daily hospitalization costs between $2000 and $5000, depending on gestational age [3]. A prolonged hospital course secondary to *S. aureus* infection can thus exponentiate costs for both families and hospital systems. A 2010 study performed in Washington, D.C. demonstrated a 40 days increase in NICU hospitalization for infants with MRSA infection, associated with a greater than $160,000 increase in cost incurred by the family [4].

MRSA colonization is a demonstrated risk factor for MRSA infection [5]. According to a recent meta-analysis, MRSA-colonized neonates were 24 times more likely to develop an MRSA infection than noncolonized neonates [6]. The prevalence of MRSA nasal colonization in neonates varies, typically between 2% and 4% [6–9], increasing to as high as 8% during an MRSA outbreak investigation [10]. Infants in the NICU are at risk for MRSA colonization secondary to transmission from colonized parents, healthcare workers, and contaminated hospital surfaces [2]. Factors associated with MRSA colonization in the NICU include lower birth weight, younger gestational age, invasive procedures, and indwelling catheters [2]. In an effort to prevent infection, decolonization with topical antimicrobials (e.g., mupirocin and chlorhexidine) is frequently employed for colonized neonates [11, 12].

Neonates colonized with MRSA are at greater risk of MRSA infection than those who are not colonized [13–15]. However, only a subset of colonized infants develop
symptomatic infection. The objective of this study was to identify factors associated with, and longitudinal outcomes related to, development of symptomatic MRSA infection in MRSA-colonized infants in the NICU.

Methods

Setting

This case-control study employed data from patients hospitalized between July 2009 and June 2018 in the St. Louis Children’s Hospital (SLCH) NICU, a level IV NICU adjacent to the Labor and Delivery Unit at Barnes-Jewish Hospital. The SLCH NICU has 85 beds (increased to 125 beds in February 2018) and ≈1300 admissions per year.

An MRSA active surveillance program was implemented in the SLCH NICU in 2004. All neonates admitted to the SLCH NICU were screened for MRSA colonization (via anterior nasal swab) at admission and weekly throughout their NICU hospitalization. Infants with positive MRSA surveillance cultures were placed in contact isolation, requiring healthcare workers to wear gowns and gloves when handling the infants for the remainder of their hospitalization. A standard decolonization protocol was introduced in 2006 for MRSA-colonized neonates consisting of intranasal 2% mupirocin ointment applied twice daily for 7 days and, at >30 weeks of gestation, a one-time bath from the neck down with 2% chlorhexidine gluconate cloths.

Study population

Infants were selected from a hospital-generated database of MRSA-colonized infants detected through active surveillance. MRSA-colonized infants who developed symptomatic infection (cases; “infected”) were matched 1:3 to MRSA-colonized infants with overlapping NICU courses who did not develop infection (controls; “colonized-only”).

Between July 2009 and June 2018, there were 7501 admissions to the SLCH NICU. Over that time, 325 infants were colonized with MRSA (4%) (Fig. 1); 62 patients (19% of those colonized) developed symptomatic MRSA infection. Infants with exposure outside of a NICU setting were excluded from the study, including infants who had been discharged from a hospital to home prior to admission to SLCH NICU (n = 1) and those hospitalized in another SLCH unit prior to NICU transfer (n = 4). Infants who spent >1 month at an outside hospital prior to SLCH NICU admission (n = 8) and infants for whom MRSA colonization was first detected >1 week after development of symptomatic infection (n = 7) were also excluded.

Symptomatic MRSA infection was defined as a bodily fluid culture-positive for MRSA (e.g., bacteremia, skin/soft tissue infection, urinary tract infection, and conjunctivitis). Patients with tracheitis, defined as a tracheal aspirate culture yielding MRSA in the presence of polymorphonuclear leukocytes on Gram stain, who required escalating respiratory support and/or demonstrated other signs of clinical instability and were treated with antibiotics, were also categorized as having symptomatic infection [16].

The study included three sets of twins (in one set, both twins were infected cases; in two sets, one twin was an infected case and the other twin was a colonized-only control) and one set of quadruplets (one infected case, three colonized-only controls). Data were abstracted for 162 mothers of the enrolled neonates. This study was approved by the Washington University School of Medicine Institutional Review Board with waiver of consent for the infants and mothers.

Data collection

Data were abstracted from electronic medical records to obtain neonates’ demographic data and characteristics of hospital stay, including sex, race, gestational age, birth weight, birth location (born at our medical center [inborn]
vs. born at an outside institution [outborn]), mode of delivery, presence of multiple gestations, comorbidities, surgical procedures, nutrition via nasogastric or orogastric tube, endotracheal intubation, number of ventilator days, presence of a central line and number of central line days, type of antibiotics received and number of systemic antibiotic days, number of systemic steroid days, exposure to maternal or donor breast milk, results of all MRSA surveillance cultures, results of all bodily fluid cultures, and receipt of intranasal mupirocin for decolonization following detection of MRSA colonization. Documentation of chlorhexidine gluconate baths was not readily available for this cohort. Maternal data obtained included race, previous MRSA-positive cultures, and presence of antepartum antibiotics.

Post-discharge longitudinal data were also obtained for neonates, when available. Information obtained included follow-up visits in newborn medicine clinic, medical diagnoses carried, subsequent hospitalizations, and subsequent surveillance and bodily fluid cultures obtained.

Statistical analyses

Statistical analyses were performed with SPSS for Windows v26 (IBM SPSS, Chicago, IL). For infected infants, data collected and presented represent characteristics of NICU course prior to infection to determine factors associated with developing an infection; these characteristics included mupirocin for decolonization, surgical procedures, nasogastric or orogastric tube, maternal breast milk, donor breast milk, intubation, ventilator days, central venous access, central line days, antibiotics, antibiotic days, inotropes, steroids, and steroid days. MRSA-colonized neonates who did and did not develop infection were compared using Mann–Whitney U test (continuous data) and Fisher’s exact test (categorical data). All tests of significance were two-tailed and p values <0.05 were considered statistically significant. To determine the factors independently associated with development of symptomatic infection, all variables tested in univariate analysis were then evaluated in a backward stepwise multivariable logistic regression. Longitudinal data analysis compared the likelihood of readmission between infected and colonized-only infants. The factors associated with hospital readmission after NICU discharge were also assessed using multivariable logistic regression; in this model, the characteristics of entire NICU course, not just prior to infection, were included.

Results

The cohort was comprised of 168 neonates: 42 MRSA-colonized infants who developed symptomatic MRSA infection (cases) and 126 MRSA-colonized infants who did not develop infection (controls). The 42 infected infants most commonly experienced tracheitis (n = 22), followed by infection of multiple sites (bacteremia plus tracheitis [n = 6], bacteremia plus skin abscess [n = 1]), bacteremia (n = 6), skin and soft tissue infection (n = 5), conjunctivitis (n = 1), and UTI (n = 1). Median time from admission to detection of colonization was 18 days (interquartile range [IQR] 10–30 days); median time from admission to infection was 24 days (IQR 14–43 days). Median time from detection of colonization to infection was 4 days (IQR 1–11 days).

Factors associated with MRSA infection

Infected infants had a lower median gestational age of 26 weeks, 1 day (IQR 24 weeks, 3 days–32 weeks, 6 days) than the colonized-only infants, with median gestational age 29 weeks, 6 days (IQR 27 weeks, 5 days–32 weeks, 5 days, p = 0.002, Table 1). Concordantly, infected infants had a lower median birth weight of 915 g (IQR 725 g–1668 g) than colonized-only infants, with median birth weight 1320 g (IQR 925 g–1913 g, p = 0.004). The proportions of male infants (30 [71%] infected, 76 [60%] colonized-only) and inborn infants (28 [67%] infected, 90 [71%] colonized-only) were similar between groups. Median time for detection of colonization (15 days for infected infants, 19 days for colonized-only infants) was also similar.

The proportion of intubated neonates was similar between groups (Table 1); however, infected infants were exposed to longer courses of mechanical ventilation (median 15 days [IQR 4–25 days]) than colonized-only infants (median 2 days [IQR 1–10 days], p < 0.001). Most infants in both groups required central vascular access (e.g., umbilical catheters, PICC lines), though infected infants had a greater number of central line days (median 16 days [IQR 12–27 days]) than colonized-only infants (median 13 days [IQR 7–21 days], p = 0.009).

Most infants in both groups (40 [95%] infected, 123 [98%] colonized-only) required a nasogastric or orogastric tube (Table 1). Thirty-four (81%) infected infants received maternal breast milk compared to 114 (91%) colonized-only infants (p = 0.11); infected infants were more likely to receive donor breast milk (18 [43%]) than colonized-only infants (18 [14%], p < 0.001). Infected infants (19 [45%]) were less likely to have received mupirocin for decolonization following detection of colonization than colonized-only infants (115 [91%], p < 0.001).

Prior to developing infection, 37 (88%) infected infants received antibiotics (16 [38%] received antibiotics with activity against MRSA); 111 (88%) colonized-only infants received antibiotics for sepsis evaluation or non-MRSA infection (50 [40%] with activity against MRSA (Table 1).
The number of antibiotic days was similar between groups: median 7 days for infected infants (for the time period prior to infection; IQR 3–13 days) and 5 days for colonized-only infants (over their entire NICU course; IQR 3–11 days, p = 0.26). All infected infants then received systemic antibiotics to treat their MRSA infection. Thirteen (31%) infected infants received inotropes prior to developing infection compared to 23 (18%) colonized-only infants (p = 0.09); 17 (41%) infected infants received systemic steroids prior to developing infection compared to 31 (25%) colonized-only infants (p = 0.08).

In multivariable analysis, infants who developed symptomatic MRSA infection were less likely to have received mupirocin for decolonization (adjusted odds ratio [aOR] 0.06, 95% confidence interval [CI] 0.02–0.19), became colonized earlier in their NICU stay (aOR 0.97, 95% CI 0.94–0.99), and received fewer days of systemic antibiotics (aOR 0.91, 95% CI 0.85–0.98) than infants who did not develop infection. Infected infants were more likely to have received donor breast milk (aOR 7.73, 95% CI 2.53–23.68) and had a more prolonged course of mechanical ventilation (aOR 1.09, 95% CI 1.04–1.14) than infants who did not develop infection.

Morbidities in patients with MRSA infection

Overall, MRSA-infected infants were exposed to longer courses of antibiotics and steroids during their hospitalization compared to colonized-only infants. Over the entire NICU course, infected infants received a median of 27 days (IQR
16–46) of systemic antibiotics and 33 days (of 29 infants that received steroids; IQR 7–68 days) of systemic steroids. Of 42 MRSA-infected infants, 14 (33%) required at least one surgical procedure prior to their infection, increasing to 25 (60%) over their entire NICU course. MRSA-infected infants had longer NICU hospitalizations (median 110 days, IQR 601–153 days) than colonized-only infants (median 60 days, IQR 40–99 days, \( p < 0.001 \)). Twenty-three (55%) infected infants were colonized at discharge compared to 55 (44%) colonized-only infants (\( p = 0.22 \)).

### Maternal data

Of 162 mothers overall, the majority (138 [85%]) were documented to have received antibiotics while admitted prior to childbirth; 12 (7%) did not receive antibiotics and 12 (7%) have unknown antibiotic use. Nine mothers received antibiotics with activity against MRSA; of these mothers, 1 (11%) of their infants developed symptomatic MRSA infection. Twenty-two mothers (14%) had a documented history of MRSA infection; 5 (23%) of these mothers were screened for MRSA colonization upon admission for childbirth and 2 (40%) were MRSA-colonized. Of these 22 mothers, 5 (23%) of their infants developed symptomatic MRSA infection.

### Longitudinal data

From July 2009 to August 2019 (the end of our data collection period), 46 patients (36%) were readmitted to SLCH within 1 year after discharge (Table 2). MRSA-infected infants (18 [53%]) were more likely to be rehospitalized within 1 year than colonized-only infants (28 [30%]; \( p = 0.02 \)). Median days from discharge to readmission did not differ significantly between groups (109 days [IQR 40–239] for MRSA-infected infants, 69 days [IQR 35–167] for colonized-only infants; \( p = 0.59 \)). One MRSA-infected patient was subsequently admitted for an MRSA abscess; two colonized-only controls (for whom longitudinal data were available) developed MRSA abscesses that did not require hospital admission.

Of 129 infants with longitudinal colonization status available, 16 were colonized with MRSA during a subsequent hospital visit (8 of 34 [24%] MRSA-infected infants, 8 of 95 [8%] colonized-only infants; \( p = 0.03 \)). Of 16 infants MRSA-colonized longitudinally, 11 (69%) were colonized at the time of discharge from their initial hospitalization; 45 (40%) infants not colonized during follow-up were colonized at the initial discharge (\( p = 0.03 \)). Patients colonized with MRSA longitudinally had longer initial NICU courses (median 124 days, IQR 58–158 days) compared to those who were not colonized longitudinally (median 82 days, IQR 48–115 days, \( p = 0.04 \)).

### Discussion

This case-control study informs factors associated with progression of MRSA nasal colonization to development of symptomatic infection in critically-ill neonates. Colonized infants who developed infection became colonized earlier in their NICU course, received fewer days of systemic antibiotics, were less likely to receive mupirocin for

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### Table 2 Subsequent admission diagnoses.

| Category                      | Specific diagnosis                                                                 |
|-------------------------------|-----------------------------------------------------------------------------------|
| **Respiratory diagnoses**     | Asthma exacerbation                                                              |
|                               | Pneumonia                                                                         |
|                               | Bronchiolitis                                                                     |
|                               | Croup                                                                             |
|                               | Aspiration pneumonia                                                             |
|                               | Cystic fibrosis exacerbation                                                     |
|                               | Subglottic stenosis after procedure                                               |
| **Cardiovascular diagnoses**  | Arrest after tracheostomy decannulation                                           |
|                               | Dehydration                                                                       |
|                               | Subglottic stenosis                                                              |
| **Infectious diagnoses**      | Abscess of left buttocks                                                         |
|                               | Pyelonephritis, renal abscess                                                    |
| **Neurologic diagnoses**      | Evaluation with EEG                                                              |
|                               | Febrile seizure                                                                   |
| **GI diagnoses**              | Bloody stools                                                                     |
| **Surgical diagnoses**        | Foreign body removal                                                             |
|                               | VP shunt malfunction                                                              |
|                               | Hiatal hernia repair                                                              |
|                               | Cleft palate repair                                                               |
|                               | Laryngotraheal reconstruction                                                     |
|                               | Peritoneal dialysis catheter placement                                            |
|                               | Dislocated left hip                                                              |
|                               | Inguinal hernia repair                                                            |
|                               | Cochlear implant placement                                                        |
|                               | Closure of ostomy and mucous fistula                                              |
|                               | Burns to bilateral feet                                                           |
| **Other diagnoses**           | Failure to thrive                                                                 |
|                               | ALTE (apparent life threatening event)                                            |
|                               | Near drowning                                                                      |

In multivariable analysis, patients who required subsequent hospitalization were more likely to have been an MRSA-infected case than a colonized-only control (aOR 2.85, 95% CI 1.29–6.30) and to have had a longer NICU course (aOR 1.01, 95% CI 1.00–1.02) during the index hospitalization. Rehospitalized infants were less likely to have received maternal breast milk (aOR 0.22, 95% CI 0.07–0.75) during the index hospitalization.
Infants who developed symptomatic infection were less likely to receive the standard decolonization protocol of 2% intranasal mupirocin twice daily for 7 days following detection of colonization. Decolonization of neonates in the NICU with mupirocin has minimal adverse effects and is efficacious in decolonizing MRSA-colonized infants [11]. Parental decolonization is another strategy that has been recently evaluated in an attempt to limit neonatal MRSA acquisition and subsequent infection. A randomized controlled trial conducted in a NICU in Baltimore demonstrated that decolonization of S. aureus-colonized parents resulted in a 57% reduction in incidence of infant acquisition of a S. aureus strain concordant with a parental strain [20, 21]. To reduce infant colonization and infection, a combination of strict adherence to infant decolonization protocols and decolonization of parents may be necessary. Mothers hospitalized in the antepartum unit are another group for whom MRSA surveillance should be considered; detecting and decolonizing these mothers prior to delivery could interrupt subsequent transmission to premature infants.

Development of symptomatic MRSA infection was associated with long-term implications during the remainder of the NICU course and after discharge. Infants who developed infection were exposed to longer courses of both systemic antibiotics and systemic steroids during their overall hospitalization than colonized-only infants, altering the already-fragile infant immune system and gut microbiome [22, 23]. Infected infants were also more likely to undergo at least one surgical procedure during their NICU hospitalization, particularly following their MRSA infection. This is likely attributable to the increase in the severity of clinical illness that results from symptomatic infection. As a result, infected infants also had a longer median length of NICU hospitalization (110 days) than colonized-only infants (60 days). After discharge, infected infants were more likely to have subsequent hospitalizations than colonized-only infants. Most rehospitalizations were secondary to respiratory distress; only one rehospitalization was secondary to reinfection with MRSA (buttock abscess).

This study has several limitations. This was a retrospective study conducted at one tertiary care center and thus the findings may not be generalizable to other geographic areas. During the study period, the SLCH NICU conducted active surveillance for MRSA only; however, the majority of symptomatic S. aureus infections were methicillin-susceptible S. aureus, limiting sample size of symptomatic infection with MRSA. At last, only a portion of the cohort had follow-up documentation available, and some infants may have been admitted to other hospitals after SLCH discharge; therefore, the longitudinal data may not be a complete reflection of colonization patterns or morbidity associated with MRSA colonization or infection in former NICU infants over time.
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**Conclusion**

In conclusion, the development of symptomatic MRSA infection in critically-ill colonized infants is multifactorial. Immature mucous membranes and skin barriers provide potential portals of entry for colonization to develop into symptomatic infection. The development of symptomatic infection is associated with increased morbidity. Reducing colonization via parental surveillance and decolonization in combination with infant decolonization may prevent future infection in already-colonized infants.

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**Author contributions**

Concept and design: CRS, PGH, PJR, MGB, BBW, and SAF. Acquisition, analysis, or interpretation of data: CRS, PGH, PJR, SH, HEW, MGB, RMT, and SAF. Drafting of the paper: CRS, PGH, PJR, SH, HEW, and SAF. Critical revision of the paper for important intellectual content: CRS, PGH, PJR, SH, HEW, MGB, RMT, BBW, and SAF. Statistical analysis: CRS, PGH, SH, HEW, and SAF. Obtained funding: SAF. Administrative, technical, or material support: MGB and RMT. Supervision: PJR, BBW, and SAF.

**Compliance with ethical standards**

**Conflict of interest**

The authors declare no competing interests.

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