The Effect of Single-Room Care Versus Open-Bay Care on the Incidence of Bacterial Nosocomial Infections in Pre-Term Neonates: A Retrospective Cohort Study

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ABSTRACT

Introduction: Nosocomial infections (NIs) are a major source of iatrogenic harm in neonatal intensive care units (NICUs). The influence of the infrastructure of NICUs on NIs is not well documented. This study aims to examine the effect of single-room units (SRU) versus open-bay units (OBU) on the incidence of NIs, including central-line-associated bloodstream infections (CLABSI), in preterm neonates.

Methods: All preterm neonates (< 32 weeks gestational age) admitted to our NICU were included. Two study periods were compared: one prior to (May 2015–May 2017) and one following (May 2017–May 2019) transition from OBU to SRU. Incidence density (number of infections per 1000 patient-days) and cumulative incidence (number of infections per 100 neonates) for NIs were calculated. CLABSIs were calculated per 1000 central-line days. U chart analysis was performed to determine special-cause variation in quarterly CLABSI and NI rates. Multivariate competing risk regression was performed to identify independent NI risk factors.

Results: Of the 712 included infants, 164 (23%) infants acquired ≥ 1 NIs. No differences were found in incidence density (13.68 vs. 12.62, p = 0.62) or cumulative incidence of NI (23.97 vs. 22.02, p = 0.59) between OBU and SRU. CLABSIs showed a similar non-significant reduction after the move (14.00 vs. 10.59, p = 0.51). U chart analysis did not identify unit transition as a potential source of special-cause variation for CLABSI and NI. Competing risks regression analysis revealed longer duration of invasive mechanical ventilation as a significant risk factor for NI (subhazards ratio: 1.03 per day on ventilation, p = 0.01).

Conclusion: Single-rooms are not associated with a significant reduction in NIs in the NICU. This study therefore does not add evidence that could support the transition to SRUs if based only on a large multimodal infection control strategy. Recommendations to build SRUs would require a wider justification, also taking into account other SRU benefits.

Keywords: Neonates; Nosocomial infections; Open-ward design; Single-room units
**Key Summary Points**

**Why carry out this study?**

Nosocomial infections (NIs) continue to be a major source of iatrogenic harm in neonatal intensive care units (NICUs) worldwide.

A recommended strategy to interrupt the transmission of pathogenic organisms and prevent NIs is to hospitalise infants in single-patient rooms; however, evidence regarding the influence of the infrastructure of NICUs on NIs remains inconclusive.

This study aimed to examine the effect of single-room units (SRU) versus open-bay units (OBU) on the incidence of NIs, including central-line-associated bloodstream infections, and overall antibiotic use in preterm neonates.

**What was learned from the study?**

SRUs are not associated with significant reduction in NIs in the NICU.

This study therefore does not provide sufficient support for single-room care as an infection-control strategy. Further investigation regarding the influence of NICU room-privatisation on the burden of NIs is needed.

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**INTRODUCTION**

Even though advances in neonatal care have led to improved survival of premature infants, nosocomial infections (NIs), also known as hospital-acquired infections, continue to be a major problem in neonatal intensive care units (NICU) worldwide [1, 2]. Prospective surveillance data over 13 non-consecutive years from the Neonatal Research Network of the Eunice Kennedy Shriver National Institute of Child Health and Human Development (NICHD) demonstrated that 21–25% of very-low-birthweight (VLBW, ≤ 1500 g) infants were found to have at least one episode of culture-confirmed sepsis, with the vast majority of infections being caused by Gram-positive organisms [1–4]. Central-line-associated bloodstream infections (CLABSIs) are the most common type of NI in neonates admitted to the NICU, and substantially factor into late-mortality, immediate- and long-term morbidity, length of hospital stay and health-care costs [5, 6]. Efforts in tackling neonatal NIs, including an ongoing assessment of associated morbidities, are required to eradicate this form of avoidable iatrogenic harm.

A recommended strategy to interrupt the transmission of pathogenic organisms and prevent NIs is to hospitalise patients in single-patient rooms, facilitating better isolation from hospital-borne infectious agents and proper execution of standard infection-control measures [7, 8]. Unit design features that are believed to contribute to a potential reduction in hospital-acquired infections include the strategic placement of hand rub dispensers closer to the point of care (i.e. near the entrance door and adjacent to the incubator), reduced physical proximity between patients (functioning as an inherent form of isolation) and limited environmental contamination as a result of reduced traffic and medical equipment requiring routine cleaning [8, 9]. However, evidence regarding the association between unit architectural design and prevention of NIs remains inconclusive. A retrospective cohort study investigating the association between NICU room configuration and colonisation by methicillin-resistant *Staphylococcus aureus* and
development of late-onset sepsis found no effect on either, even after controlling for demographic, clinical and unit-based characteristics [9]. In contrast, a recent systematic review evaluating the difference between single-room care versus open-bay care on a variety of neonatal outcomes in preterm infants did report a reduced incidence of sepsis in infants hospitalised in single rooms [10].

Following transition from an open-bay unit (OBU) to a newly constructed single-room unit (SRU) NICU at Leiden University Medical Center (LUMC) on May 15, 2017, we examined the effect of changing from an open, multiple-bed ward to an entirely single-room ward on the incidence of nosocomial infections, including CLABSI, and overall antibiotic use.

METHODS

Setting and Study Design

The LUMC is a tertiary university-affiliated hospital with a 25-bed level-III NICU constructed as a single-room facility with an average annual admission rate of approximately 500–600 in- and out-born neonates. The ward contains 16 single-patient rooms and four twin-rooms (Supplementary Figure S1). In addition, a separate negative-pressure isolation room designed to prevent the spread of airborne pathogens is located adjacent to the main entrance of the ward. Each infant is hospitalised in a private room equipped with 1 hand-washing facility with touch taps and a dispenser with alcohol-based hand rub. Additional hand rub dispensers are available adjacent to each incubator. The average nurse-to-patient ratio is 1:1 for infants with high-dependency care, and 1:2 for infants requiring intermediate care. Prior to the transition to the new single-room facility, the NICU consisted of three open-bay rooms, two which served as intensive care (IC) units and one as a high-care (HC) unit. The two IC units were configured with, respectively, nine and seven beds, and the HC unit with nine beds. Bed occupancy rates depended on volatility in demand and staffing levels. Each unit was equipped with one sink (Supplementary Figure S2). Hand rub dispensers were located next to each incubator and at the central nursing station. Depending on the level of care required, the standard nurse-to-patient ratio was 1:2 and 1:3 in the IC and HC units, respectively. The average area surrounding each incubator was approximately 2 m².

We conducted a retrospective cohort study to evaluate the rate of nosocomial infections and antibiotic consumption in premature infants before and after transition to the new single-room NICU. Two study periods of 2 years each were compared: one prior to (May 15, 2015–May 14, 2017) and one following (May 16, 2017–May 15, 2019) unit transition. Approval and a waiver from the need to provide written informed consent were obtained from the institutional review board of LUMC (G19.070).

Study Subjects and Data Collection Procedures

Neonates born at < 32 weeks gestational age and admitted to the NICU between May 15, 2015 and May 15, 2019 were included in this study. Infants born on the day of the unit transition (May 15, 2017) and those who were admitted to both unit types were excluded. Only data from first admissions were analysed.

For all study participants, the following data were extracted from electronic medical records for the period between admission and hospital discharge: birth date, gestational age, birth weight, sex, delivery mode (i.e. caesarian section or vaginal delivery), 5-min Apgar score, presence of major congenital anomalies, presence of surgical pathology, exposure to invasive mechanical ventilation, duration of invasive mechanical ventilation, exposure to a central-line, exposure to intravenous antibiotic therapy ≤ 24 h postpartum and ≥ 72 h after admission, total length of antibiotic treatment, prophylactic fluconazole use, length of hospital stay, in-hospital neonatal mortality and infection-related mortality (i.e. death attributed to infection as either the immediate or underlying cause). Information on NIs was obtained by reviewing patients’ medical records, including laboratory
(i.e. C-reactive protein), microbiology (i.e. organisms isolated from clinical cultures) and radiology (i.e. ultrasound, chest and abdominal radiography) results. Data pertaining to CLABSI included central-line type (i.e. umbilical venous, umbilical arterial or central-venous catheter), number of central-lines placed per infant, total duration of central-line use and isolated pathogenic microorganism(s) in case of a confirmed CLABSI.

**Sepsis Work-Up and Culture Collection**

Evaluation for sepsis was performed at the discretion of the attending physician in accordance with hospital protocol and included blood-culture analysis and at least one quantitative C-reactive protein (CRP) measurement 18 h after the initial sepsis work-up. Only culture results for specimens collected ≥ 72 h after NICU admission were considered. In case an infant was evaluated for sepsis more than once, all new unrelated episodes – i.e. absence of clinical signs or positive cultures between two episodes – were regarded as separate infection episodes.

**Definitions**

Nosocomial infection was defined as isolation of a pathogenic organism from a blood or other sterile fluid culture obtained ≥ 72 h after admission in combination with clinical symptoms of infection (temperature instability (< 36.5 or ≥ 38°C), abdominal distension, feeding intolerance, respiratory distress, increase in ventilation requirements, apnea, hypotension, hyper-/hypoglycemia, metabolic acidosis, abnormal laboratory signs (i.e. leukocytosis, thrombocytopenia) or a CRP ≥ 10 mg/L). CLABSI was defined as a laboratory-confirmed bloodstream infection occurring ≥ 72 h after birth and associated with an indwelling central-line and not related to an infection elsewhere. In addition, the definition requires that the central-line had been in place for > 2 calendar days, up until the day of removal or the following day. Confirmed CLABSI were subsequently categorised into one of three criteria based on the isolated pathogen (Supplementary Table S1). A positive culture with coagulase-negative staphylococci (CoNS) was considered an infection when a CRP ≥ 10 mg/L was measured within 36 h of infection onset. Cultures yielding other low virulence organisms (*Corynebacterium* spp., *Micrococcus* spp.) without elevated inflammatory markers were regarded as contamination and removed from further analyses. Given their contribution to short- and long-term neonatal morbidity, and in order to provide a comprehensive measure of the burden of NIs among preterm neonates, additional non-CLABSI NI types were also evaluated. As standardised definitions for NIs within the neonatal population are lacking, definitions for the non-CLABSI NI types were developed using well-established reference sources and adjusted according to specific neonatal symptomatology and diagnostic procedures. The diagnosis of a NEC was based exclusively on clinical parameters, as data on the prevalence of concurrent positive blood cultures in infants diagnosed with NEC are inconclusive [11]. Only grades IIA–IIIB according to Bell’s staging criteria were collected [12]. Remaining definitions pertaining to each NI type are listed in Supplementary Tables S1 and S2.

**Statistical Analysis**

Data are reported as mean and standard deviation (SD), median and interquartile range (IQR) or absolute number and percentages, as appropriate. In univariate analysis, categorical data were analysed using the Chi-square test or Fisher’s exact test, and continuous data using the independent *t* test or Mann–Whitney *U* test, as appropriate. Rates for NIs were calculated as the number of infection episodes per 1000 patient-days (incidence density) and number of infection episodes per 100 infants (cumulative incidence rate). CLABSI rates were calculated per 1000 central-line days. Considering infections as rare events, differences in infection rates in Poisson distribution were calculated using the exact conditional test. Shewhart *U* charts were used to evaluate the presence of special-cause variation with regards to quarterly
CLABSI and NI rates [13]. To evaluate change in antibiotic use between the two study periods, total days of treatment (DOT) per antibiotic agent and length of treatment (LOT) were calculated. If an infant received two or three different antibiotic agents within 1 day, each was counted as a separate DOT. Finally, to identify predictors for NI, univariate and multivariate Fine-Gray Competing Risk Regression models were created in which death without infection was modelled as a single competing outcome. Given that the outcome of death may modify the probability that an infection occurs, the competing risk modelling approach was chosen to allow for direct assessment of the association between a covariate and the cumulative incidence of infection. Unit type was included as the sole variable in the univariate model as the main outcome of interest. For the multivariate model, predictors were chosen based on clinical relevance. Retainment in or elimination from the model was based on maximum log-likelihood estimates using the stepwise selection approach. Subdistribution hazard (sDH) ratios comparing the probability of NI for each predictor value, conditional upon survival or occurrence of the competing risk (death), were computed along with corresponding 95% confidence intervals (CI). All p values were two-tailed and p < 0.05 was regarded as statistically significant. All statistical analyses were performed using R (v.3.6.2) [14].

RESULTS

Demographic Characteristics

Derivation of the study sample is shown in Fig. 1. Totals of 367 and 345 premature infants were admitted to the open-bay and single-room units, respectively, whose demographic characteristics are described in Table 1. There were significantly more infants in whom a central-line was inserted (61% vs. 48%, p = 0.005) during the open-bay period compared to after the conversion to single-room units. No other significant differences in neonatal characteristics between the two study periods were identified.

Incidence of Nosocomial Infections

A total of 200 NI episodes occurred in 164 (23%) infants (Table 2). No differences were found in the incidence density per 1000 patient-days (13.68 vs. 12.62 p = 0.62) nor cumulative incidence rate per 100 neonates (23.97 vs. 22.02, p = 0.59) between the historic open-ward and new single-room unit.

Apart from a slight increase in the number of primary blood stream infections and upper respiratory tract infections, an overall decrease was seen in the number of episodes for the remaining infection types after transition to the new unit, despite not being statistically significant (Table 3). No infant developed a meningitis during either study period.

During both study periods, three neonates died of a hospital-acquired infection, with mortality directly attributable to infection being 0.4% (Table 1). Of these three neonates, two were hospitalised in the new SRU. The cause of death of these three neonates were, respectively, E. coli sepsis-related multiorgan failure, methicillin-sensitive S. aureus-related respiratory and circulatory failure, and fulminant sepsis after NEC.

Figure 2 details the quarterly NI rates across the entire 4-year study period. The mean NI rate was 11.2 per 1000 patient-days. U chart analysis did not reveal any occasion in which the variation in NI incidence was higher or lower than expected by random (common-cause) variation, particularly around the time of unit transition.

Central-line Characteristics and Incidence of CLABSI

As the most common type of NIs in neonates, we evaluated the effect of unit transition on the occurrence of CLABSI by assessing changes in central-line use, dwell-time and incidence of central-line infections among neonates with one or more central-lines (Table 4). Over the two study periods, 779 central-lines with 2963 central-line days were placed in 390 infants, with significantly more central-lines placed during the open-bay period compared to the single-room period (430 vs. 349, p = 0.004). No
difference was found in average number of central-line days per child, type of inserted central-line, and average age at central-line insertion and removal. Overall, 23 neonates developed at least one CLABSI in the OBU compared to 14 in the SRU, although this difference was not statistically significant ($p = 0.51$). The 25% reduction in CLABSI rate (14–10.59 per 1,000 central-line days) after unit transition was also not statistically significant ($p = 0.51$).

Figure 3 shows the quarterly CLABSI rates across the entire 4-year study period, with the mean CLABSI rate being 12.5 infections per 1000 line-days. No special-cause signalling, specifically around unit transition, was identified by $U$ chart analysis. A steep increase in the infection rate in the period May–July 2019 is present, likely resulting from a lower number of line-days (36.37 line-days) compared to prior quarters.

**Pathogen Distribution**

Supplementary Table S3 details the pathogen distribution of the most prevalent NI types, namely, CLABSI and primary bloodstream infection (PBSI). For both infection types, the majority of monomicrobial infections were caused by Gram-positive cocci (CLABSI: 92%, PBSI: 84%), with Gram-negative organisms accounting for 5.4 and 7% of CLABSI and PBSI episodes, respectively. There was a slight decrease in CLABSIs and PBSIs owing to CoNS after unit transition (CLABSI: 87% vs. 71%, PBSI: 70% vs. 68%). Other frequently isolated Gram-positive organisms included $S. aureus$ and

**Fig. 1** Flow chart of study participant enrollment
### Table 1 Infant and maternal characteristics

| Characteristic                        | OBU (\(n = 367\)) | SRU (\(n = 345\)) | \(P\) value |
|---------------------------------------|--------------------|--------------------|-------------|
| Sex (female), \(n\) (%)              | 171 (47)           | 175 (46)           | 0.30        |
| Birth weight (g), mean (SD)           | 1285 (377)         | 1240 (382)         | 0.11        |
| Number of VLBW infants, \(n\) (%)     | 260 (71)           | 253 (73)           | 0.51        |
| Number of ELBW infants, \(n\) (%)     | 91 (25)            | 99 (29)            | 0.28        |
| Gestational age (weeks), median (IQR)| 29 (4)             | 29 (4)             | 0.84        |
| Delivery by caesarian section, \(n\) (%)| 196 (53)           | 185 (54)           | 1           |
| 5-min Apgar score, median (IQR)\(^a\) | 8 (2)              | 8 (2)              | 0.17        |
| Antenatal steroids, \(n\) (%)\(^b\)  | 340 (93)           | 321 (93)           | 0.95        |
| Postnatal steroids, \(n\) (%)        | 19 (5.2)           | 20 (5.8)           | 0.84        |
| Exposure to central-line, \(n\) (%)  | 223 (61)           | 167 (48)           | 0.005       |
| Major congenital anomaly, \(n\) (%)  | 10 (2.7)           | 3 (0.9)            | 0.12        |
| Exposure to invasive mechanical ventilation, \(n\) (%)\(^d\) | 140 (38) | 129 (37) | 0.89 |
| Total ventilation-days, \(n\)        | 996                | 956                | 0.97        |
| Duration of invasive mechanical ventilation per infant (days), median (IQR)\(^e\) | 4 (7) | 5 (6) | 0.97 |
| Surgical pathology, \(n\) (%)        | 15 (4.1)           | 11 (3.2)           | 0.66        |
| Prophylactic fluconazole use, \(n\) (%) | 57 (16)           | 62 (18)           | 0.44        |

Significant \(P\) value shown in bold

\(OBU\) open-bay unit, \(SRU\) single-room unit, \(SD\) standard deviation, \(gr\) grams, \(VLBW\) very-low birth-weight, \(ELBW\) extremely-low birth-weight, \(IQR\) interquartile range

\(^a\) Results based on 365 and 342 infants in OBU and SRU, respectively (five missing variables)

\(^b\) Results based on 361 and 341 infants in OBU and SRU, respectively (ten missing variables)

\(^c\) Defined as a condition that leads to significant medical, social or cosmetic consequences and typically requires medical intervention such as oral facial clefts, cyanotic heart defects, neural tube defects and limb deficiencies

\(^d\) Ventilation modalities include conventional mechanical ventilation with pressure support or high frequency oscillation (HFO)

\(^e\) Results displayed for infants exposed to invasive mechanical ventilation only

### Table 1 continued

| Characteristic                        | OBU (\(n = 367\)) | SRU (\(n = 345\)) | \(P\) value |
|---------------------------------------|--------------------|--------------------|-------------|
| Total patient-days, \(n\)             | 8185               | 6969               | 0.36        |
| Length of hospital stay per child (days), median (IQR) | 13 (25) | 13 (23) | 0.36 |
| In-hospital mortality, \(n\) (%)      | 15 (4.1)           | 20 (5.8)           | 0.38        |
| Infection-related mortality, \(n\) (%)| 1 (0.3)            | 2 (0.6)            | 0.96        |

### Table 2 Rates of nosocomial infections

| Characteristic                        | OBU   | SRU   | \(P\) value |
|---------------------------------------|-------|-------|-------------|
| Total infection episodes, \(n\)      | 112   | 88    | –           |
| Incidence density per 1000 patient-days | 13.68 | 12.62 | 0.62       |
| Infants with an infection, \(n\)     | 88    | 76    | –           |
| Cumulative incidence per 100 infants  | 23.97 | 22.02 | 0.59        |

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Enterococcus spp. (E. faecium), the former increasing in prevalence among PBSIs following transition to the SRU. Two PBSI episodes were found to be polymicrobial, both predominated by Gram-negative organisms. Only a small proportion of CLABSI and PBSI episodes were caused by Gram-negative organisms and fungi throughout the entire study period.

Antibiotic Exposure

No difference between the two units was found in number of neonates who received antibiotic treatment within the first 24 h postpartum (62% vs. 57%, \( p = 0.19 \)) (Supplementary Table S4). Likewise, no difference between the units was found in the proportion of neonates who were treated with one or more courses of antibiotics \( \geq 72 \) h after admission (64% vs. 57%, \( p = 0.06 \)). Total LOTs of 2832 and 2413 days were registered in the OBU and SRU, respectively, with the normalised difference not statistically significant (345 vs. 346 per 1000 patient-days, \( p = 0.51 \)). Similarly, no significant decrease was found in antibiotic consumption accounting for each agent (DOT, 616 vs. 607 per 1,000 patient-days, \( p = 0.99 \)).

Multivariate Analysis by Fine-Gray Competing Risk Regression Model

Supplementary Table S5 shows the subdistribution hazard ratios estimated from the Fine–Gray competing risk regression model. In univariate analysis, unit type was not associated with a change in risk of infection (sDH: 0.95, \( p = 0.78 \)). In multivariate analysis, 12 variables, namely unit type, gestational age, sex, 5-min Apgar score, birth weight, birth modus, antenatal steroids, duration of invasive mechanical ventilation, exposure to a central-line, central-line dwell-time, antibiotic treatment within 24 h after birth and length of hospital stay were selected to be evaluated as potential risk factors for NI. Only duration of invasive mechanical ventilation was identified as a minor risk factor (sDH: 1.03, \( p = 0.01 \)). No other variables, including single-room care, were significantly associated with NI in multivariate analysis.

DISCUSSION

We conducted a retrospective cohort study to assess the incidence of hospital-acquired infections in premature neonates admitted to a newly designed ward with single-rooms versus a historic ward with three multi-bed wings. Our findings revealed a non-significant overall reduction in NIs after unit transition. Similar non-significant reductions were found in the CLABSI rate and the number of other NI types after unit transition. Gram-positive organisms, in particular CoNS, were the most frequently isolated pathogens during both study periods. We found no significant reduction in the proportion of infants treated with antibiotics nor in

| Table 3 Number of infection episodes per infection type |
|--------------------------|--------------------------|--------------------------|--------------------------|
| Infection type           | Unit type                | OBU | SRU | \( P \) value |
| Central-line associated bloodstream infection (CLABSI) | Central-line associated bloodstream infection, Persistent bacteremia after CLABSI, Primary bloodstream infection (PBSI), Secondary bloodstream infection (SBSI), Hospital-acquired pneumonia (HAP), Ventilator-associated pneumonia (VAP), Upper respiratory tract infection (URTI), Necrotising enterocolitis (NEC), Urinary tract infection (UTI), Skin and/or soft tissue infection, Conjunctivitis | 23  | 14 | 0.24 |
| Persistent bacteremia after CLABSI | 3  | 1 | 0.66 |
| PBSI                     | 40 | 44 | 0.52 |
| SBSI                     | 5  | 3  | 0.79 |
| HAP                      | 2  | 0  | 0.51 |
| VAP\(^a\)                | 3 (3.01) | 2 (2.09) | 1 |
| URTI                     | 1  | 4  | 0.33 |
| NEC                      | 25 | 17 | 0.36 |
| UTI                      | 3  | 2  | 1 |
| Skin and/or soft tissue infection | 3  | 1  | 0.66 |
| Conjunctivitis           | 4  | 0  | 0.15 |
| Total                    | 112 | 88 | 0.15 |

\( CLABSI \) central-line associated bloodstream infection, \( PBSI \) primary bloodstream infection, \( SBSI \) secondary bloodstream infection, \( HAP \) hospital-acquired pneumonia, \( VAP \) ventilator-associated pneumonia, \( URTI \) upper respiratory tract infection, \( NEC \) necrotising enterocolitis, \( UTI \) urinary tract infection

\(^a\) Results in parentheses represent incidence per 1000 ventilation-days

Enterococcus spp. (E. faecium), the former increasing in prevalence among PBSIs following transition to the SRU. Two PBSI episodes were found to be polymicrobial, both predominated by Gram-negative organisms. Only a small proportion of CLABSI and PBSI episodes were caused by Gram-negative organisms and fungi throughout the entire study period.
overall antibiotic consumption. Only duration of invasive mechanical ventilation, which was not different between the unit types, was identified as a risk factor for NI.

Even though the present study found a decrease in the number of infections, this decrease was statistically non-significant, and it also remains unclear if, and to what extent, the transition from OBU to SRU contributed to this decrease. While we aimed to identify unit transition as a potential source of special-cause variation in our $U$ charts, we found only random variation present, suggesting that no particular infection prevention element or change in clinical practice contributed to a statistically significant reduction in NI rates during the study period. Moreover, unwavering adherence to hand hygiene practices may have further diminished the impact of unit configuration. The overall decrease in the number of children with an infection and the number of infection episodes may also have been part of an existing declining trend rather than a result of the new SRU design.

It is important to note that our NI rates are slightly higher compared to those reported in other studies [15–18], presumably because we chose to also report less severe (i.e. skin and eye) and secondary infections (i.e. persistent bacteremia after CLABSI, secondary bloodstream infection). Although not immediately life-threatening, these infection types still act as an important reservoir of pathogenic organisms and may lead to clinically manifested consequences [19]. The substantial variability in definitions of neonatal CLABSI across studies in combination with different baseline populations render an adequate comparison of CLABSI rates difficult.

Unlike the number of CLABSI, which did not decrease significantly after conversion to the SRU, the number of central-lines placed was significantly lower. At our institution, central-lines are most frequently inserted in infants born at < 28 weeks’ gestation and/or with a

Fig. 2 $U$ chart for overall NI rates by 3-month periods. The central line (CL) represents the internal mean NI rate from May 15, 2015 to May 15, 2019. The upper and lower limits represent the upper and lower control limits, respectively (UCL, LCL). LCL approaches zero from May 2018 to November 2018 and from February 2019 onwards and is therefore not depicted. Final quarter (May–Jul 2019) displayed as full 3-month period, although data present up until May 15, 2019.
All insertion procedures are performed by experienced neonatologists, physician assistants or competent paediatric residents according to a standardised institution-specific protocol and checklist. Although no formal changes in the central-line insertion and maintenance policy were made during either study periods, the decrease in the number of central-lines placed is suggestive of a more restrictive policy regarding the use of invasive medical devices, which may eventually lead to a lower CLABSI rate. The absence of meningitis in either study period may in part be due to our stringent definition which does not presumptively rely only on abnormal CSF parameters (i.e. cell count, glucose, protein levels). Also, CoNS was the primary pathogen isolated from all CSF cultures collected from infants clinically suspected of a meningitis, suggesting specimen contamination instead of true infection.

Our non-significant reduction in NIs following unit transition contrasts with findings from a recent systematic review and meta-analysis assessing the effect of single-family rooms versus open-bay units on a variety of neonatal outcomes in preterm infants, including late-onset sepsis [10]. Nevertheless, only six studies assessing culture-proven late-onset sepsis were included, four of which had moderate risk of bias and thus a possible overestimation of the intervention effect. Additionally, compared to our study, three of the six studies included found a relatively larger reduction (5–9%), suggestive of higher pre-unit transition rates resulting in larger reductions [15,20]. Moreover, the true effect of single-room care on late-onset infection rates may have been overestimated, as publication bias was not assessed. Recommendations to build SRUs would therefore require a wider justification also taking into account other SRU benefits.

A secondary objective of our study was to examine whether single-room care would lead to a decrease in overall antibiotic exposure, perhaps as a result of a concomitant decline in the number of NIs. We found no difference in the number of infants treated with antibiotics in the first 24 h after birth between SRU and OBU. This empiric treatment is often initiated due to a suspected early-onset infection, the

| Characteristic                      | OBU      | SRU      | P value |
|------------------------------------|----------|----------|---------|
| Central-lines placed, n            | 430      | 349      | 0.004   |
| Total central-line days, n         | 1642.5   | 1321.1   | 0.24    |
| Central-line days per infant, median (IQR) | 6.71 (5.28) | 7.03 (4.02) | 0.24    |
| Number of VLBW infants with a central-line, n (%)a | 188 (72) | 155 (61) | 0.07    |
| Number of ELBW infants with a central-line, n (%)a | 87 (96) | 89 (90) | 0.88    |
| Central-line type, n (%)           |          |          |         |
| UVC                                | 173 (40) | 135 (39) | 0.19    |
| UAC                                | 114 (27) | 80 (23)  |         |
| CVC                                | 143 (33) | 134 (38) |         |
| CLABSI episodes, n b               | 23       | 14       | 0.51    |
| Infants with ≥ 1 CLABSI(s), n c    | 23       | 14       | 0.64    |
| Age at insertion (d), median (IQR) | 0.85 (0.48) | 0.92 (0.69) | 0.07    |
| Age at removal (d), median (IQR)   | 8.44 (5.68) | 8.93 (5.93) | 0.13    |
| CLABSI incidence per 1000 central-line days | 14.00 | 10.59 | 0.51 |

Significant P value shown in bold.
IQR interquartile range. VLBW very-low-birth-weight (<1500 g). ELBW extremely-low-birth-weight (<1000 g). CLABSI central-line associated bloodstream infection. UVC umbilical-venous catheter. UAC umbilical-arterial catheter. CVC central-venous catheter. d days.

a VLBW total group consists of 260 and 253 infants in OBU and SRU, respectively. ELBW total group consists of 91 and 99 infants in OBU and SRU, respectively.
b Calculated difference based on total number of central-lines placed (n = 779).
c Calculated difference based on total number of infants with a central-line (n = 390).
incidence of which is unlikely to have been impacted by change in unit configuration. In parallel with the nonsignificant decrease in the number of NI, fewer infants received antibiotic treatment \( \geq 72 \) h after admission. Nevertheless, LOT and DOT rates remained relatively stable between the two study periods, indicating that single-room care did not impact overall consumption.

**Strengths and Limitations**

The strengths of our study include the meticulous data collection and clearly defined NI types. However, our data also have several limitations. First, the limited power of our study may have inherently hindered the results from reaching statistical significance. A repeat analysis with a larger study sample and a longer time series is therefore desirable. Second, we were unable to correct for potential prevailing local temporal trends in NIs due to lack of available data. Analysis using segmented interrupted-time series with monthly incidence rates as opposed to a simpler pre- and post-analysis would have allowed for a more refined assessment of the effect of unit transition on infection rates. Lastly, we were unable to identify and model for all other potential modifications in infection-control practices that may have taken place as part of continuous quality improvement efforts.

**CONCLUSION**

Based on the data available in this study, the notion that the transition from a historic, open-bay ward to a new single-room unit provides added protection against hospital-acquired infections in preterm infants is not supported. Given the complex, multifactorial nature of these infections, improvements primarily lie in whether basic measures such as maintaining adequate staffing levels, limiting the use of

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**Fig. 3** *U* chart for CLABSI rates by 3-month periods. The central line (CL) represents the internal mean CLABSI rate from May 15, 2015 to May 15, 2019. The upper line represents the upper control limit (UCL). The lower control limit (LCL) approaches zero and is therefore not depicted. Final quarter (May–Jul 2019) displayed as full 3-month period, although data present up until May 15, 2019.
invasive medical devices, ensuring high level of competency for insertion and maintenance of central-lines, antibiotic stewardship and adherence to hand hygiene, are met. Redesigning a NICU involves substantial budgetary implications and staff adjustment and thus must be based on solid evidence. While our findings thus provide insufficient support for single-room care as an infection-control strategy, further investigation regarding the influence of NICU room-privatisation on the burden of NIs is therefore warranted, preferably with larger study samples spanning longer time series and a randomised control design.

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Data Availability The datasets generated during and/or analysed during the current study are not publicly available due to the privacy of the participants.

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