Resuscitation of Preterm Neonates With Limited Versus High Oxygen Strategy

**Objective:** To determine whether a limited oxygen strategy (LOX) versus a high oxygen strategy (HOX) during delivery room resuscitation decreases oxidative stress in preterm neonates.

**Methods:** A randomized trial of neonates of 24 to 34 weeks’ gestational age (GA) who received resuscitation was performed. LOX neonates received room air as the initial resuscitation gas, and fraction of inspired oxygen (FiO2) was adjusted by 10% every 30 seconds to achieve target preductal oxygen saturations (SpO2) as described by the 2010 Neonatal Resuscitation Program guidelines. HOX neonates received 100% O2 as preductal oxygen saturations (SpO2) as described by the 2010 Neonatal Resuscitation Program guidelines. HOX neonates received 100% O2 as initial resuscitation gas, and FiO2 was adjusted by 10% to keep preductal SpO2 at 85% to 94%. Total hydroperoxide (TH), biological antioxidant potential (BAP), and the oxidative balance ratio (BAP/TH) were analyzed in cord blood and the first hour of life. Secondary outcomes included delivery room interventions, respiratory support on NICU admission, and short-term morbidities.

**Results:** Forty-four LOX (GA: 30 ± 3 weeks; birth weight: 1678 ± 634 g) and 44 HOX (GA: 30 ± 3 weeks; birth weight: 1463 ± 606 g) neonates were included. LOX decreased integrated excess oxygen ($\sum$FiO2 \times \text{time [min]}) in the delivery room compared with HOX (401 ± 151 vs 662 ± 249; P < .01). At 1 hour of life, BAP/TH was 60% higher for LOX versus HOX neonates (13 [9–16] vs 8 [6–9]) μM/U.CARR, $P < .01$). LOX decreased ventilator days (3 [0–64] vs 8 [0–96]; $P < .05$) and reduced the incidence of bronchopulmonary dysplasia (7% vs 25%; $P < .05$).

**Conclusions:** LOX is feasible and results in less oxygen exposure, lower oxidative stress, and decreased respiratory morbidities and thus is a reasonable alternative for resuscitation of preterm neonates in the delivery room. *Pediatrics* 2013;132:e1488–e1496

**Abbreviations:**
- BAP—biological antioxidant potential
- BPD—bronchopulmonary dysplasia
- CPAP—continuous positive airway pressure
- DR—delivery room
- FiO2—fraction of inspired oxygen
- GA—gestational age
- HFOV—high-frequency oscillatory ventilation
- HOX—high oxygen strategy
- HR—heart rate
- IVH—intraventricular hemorrhage
- LOX—limited oxygen strategy
- NRP—Neonatal Resuscitation Program
- OB—obstetrical
- PPV—positive pressure ventilation
- PVL—periventricular leukomalacia
- ROP—retinopathy of prematurity

Dr Kapadia conceptualized and designed the study, recruited patients, and designed the data collection instruments; participated in collecting the data, analysis of the data, and interpretation of data; and drafted the initial manuscript, wrote the last draft of the manuscript, and approved the final manuscript as submitted. Dr Chalak participated in concept, design, analysis, and interpretation of data; reviewed and revised the manuscript, and approved the final manuscript as submitted. Dr Sparks participated in recruitment, design, analysis, and interpretation of data; reviewed and approved the final manuscript as submitted. Mr Allen participated in data, collection, and analysis of data, and reviewed and approved the final manuscript as submitted. Dr Savani participated in concept, design, analysis, and interpretation of data; and critically reviewed and approved the final manuscript as submitted. Dr Wyckoff participated in conceptualization and design of the study; supervised the recruitment, designing of data, and collection instruments; participated in analysis of the data and interpretation of the data; and critically reviewed, revised, and approved the final manuscript as submitted. This trial has been registered at www.clinicaltrials.gov (identifier NCT01687904).

(Continued on last page)
Use of high oxygen (O₂) concentrations during delivery room (DR) resuscitation generates bursts of reactive oxygen and nitrogen species that can overwhelm newborn antioxidant capacity and damage cell components such as lipids, proteins, RNA, and DNA. Even brief 100% O₂ exposure during resuscitation increases oxidative stress and reduces antioxidant capacity as measured by using biomarkers such as total hydroperoxide (TH), biological antioxidant potential (BAP), reduced glutathione/oxidized glutathione ratio, and superoxide dismutase activity. Oxidative stress increases newborn morbidities such as bronchopulmonary dysplasia (BPD), retinopathy of prematurity (ROP), intraventricular hemorrhage (IVH), periventricular leukomalacia (PVL), and mortality. Preterm neonates are especially vulnerable to O₂ toxicity because they frequently require resuscitation after birth and receive supplemental O₂ despite reduced antioxidant defenses that mature later in gestation.

In 2005, the American Academy of Pediatrics/American Heart Association Neonatal Resuscitation Program (NRP) guidelines preferentially recommended the use of 100% O₂ but allowed providers to begin resuscitation with less O₂ and recommended using pulse oximetry to adjust O₂ concentrations as needed. These guidelines emphasized that no studies justified any particular starting O₂ concentration. Subsequently, 4 small trials demonstrated that resuscitation of preterm neonates with an initial fraction of inspired oxygen (FiO₂) of 0.21 to 0.30 was feasible when FiO₂ was titrated against preductal oxygen saturations (SpO₂) and for bradycardia. However, the optimal O₂ strategy during resuscitation remains unknown.

Despite no studies comparing outcomes of neonatal resuscitation targeted at various oxyhemoglobin concentrations, the 2010 NRP guidelines recommended that preterm neonates at birth have O₂ titrated to achieve O₂ saturations in the interquartile range of healthy term newborns and that excessive O₂ should be avoided. To date, no randomized trials of strategies to achieve the NRP-recommended interquartile range of preductal saturations have been conducted in preterm neonates.

We hypothesized that a limited O₂ strategy (LOX) with resuscitation initiated with 0.21 FiO₂ and adjusted to meet NRP transitional goal saturations would increase the oxidative balance ratio (BAP/TH) by at least 15% compared with a high O₂ strategy (HOX) in which resuscitation was initiated with pure O₂ and titrated for a targeted SpO₂ of 85% to 94%.

METHODS

Patients

This study was a prospective randomized trial conducted from August 2010 to January 2011 at Parkland Hospital, Dallas, Texas. The project was approved by the institutional review board of the University of Texas Southwestern Medical Center. The 2010 NRP guidelines had not been released or implemented at study initiation, and 2005 NRP was in use. Because both LOX and HOX were consistent with the 2005 NRP guidelines and because there was equipoise regarding the 2 treatment arms, the institutional review board permitted the trial to proceed without antenatal consent as long as parental informed consent was subsequently obtained. All inborn neonates of obstetrical (OB) gestational age (GA) 24 0/7 to 34 6/7 weeks for whom the high-risk resuscitation team was present at birth and who required active resuscitation were included. Active resuscitation was defined as need for blow-by O₂ for low SpO₂, positive pressure ventilation (PPV), continuous positive airway pressure (CPAP), or ventilation via an endotracheal tube. Neonates were excluded for nonviability, prenatally diagnosed cyanotic congenital heart disease, if no active resuscitation was required, or if preductal SpO₂ could not be measured. Multiple gestations were assigned randomly according to the individual neonate.

Procedures

The high-risk neonatal resuscitation team attended delivery of all neonates <35 weeks’ OB GA. Radical oximeters (Masimo Corporation, Irvine, CA) were set to maximal sensitivity with 2-second averaging. The probe was placed on the preductal hand within the first 30 seconds of life by using previously described strategies to optimize time to reliable signal.

A noninvestigator used a permuted design in blocks of 8 to determine randomization sequence and stratified participants into OB-estimated GA groups: 24 0/7 to 28 6/7 weeks and 29 0/7 to 34 6/7 weeks. Allocation was concealed via serially numbered, sealed, opaque envelopes that were opened sequentially by the resuscitation team when the need for resuscitation was recognized. Resuscitation started with room air while the envelope was opened and then immediately changed to LOX or HOX as assigned in a nonblinded fashion.

Limited Oxygen Strategy

Resuscitation was initiated with 21% O₂ for LOX neonates. Supplemental O₂ was given for the following indications: (1) heart rate (HR) <100 beats per minute after 30 seconds of effective ventilation; or (2) lower limit of goal saturations not achieved. Targeted preductal saturations after birth were derived by approximation of interquartile values for healthy term neonates (Table 1). FiO₂ was adjusted by 10% in 30-second intervals to maintain saturations in the interquartile range. If HR remained <60 beats per minute despite 30 seconds of PPV, FiO₂ was increased to 100% until recovery of HR to >100 beats per minute.
TABLE 1  Targeted Preductal SpO2 After Birth

<table>
<thead>
<tr>
<th>Time</th>
<th>Saturation Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 min</td>
<td>80%–85%</td>
</tr>
<tr>
<td>2 min</td>
<td>65%–70%</td>
</tr>
<tr>
<td>3 min</td>
<td>70%–75%</td>
</tr>
<tr>
<td>4 min</td>
<td>75%–80%</td>
</tr>
<tr>
<td>5 min</td>
<td>80%–85%</td>
</tr>
<tr>
<td>10 min</td>
<td>85%–94%</td>
</tr>
</tbody>
</table>

High Oxygen Strategy

For HOX neonates, resuscitation was initiated with 100% O2 and adjusted every 30 seconds by 10% to meet target SpO2 of 85% to 94%.

Resuscitation Management

Apart from the randomized O2 strategy, resuscitation followed 2005 NRP guidelines. Treatment failure was defined as HR <60 beats per minute despite 30 seconds of effective PPV. If pulse oximetry did not register stable values, resuscitation continued at the current FIO2 as long as an HR >100 beats per minute was maintained.

Subsequent NICU Management

After NICU admission, all care decisions (including ventilator management) were at the discretion of the attending neonatologist. Target SpO2 was 88% to 94% throughout the NICU stay per unit policy. Infants with an O2 requirement at 36 weeks’ corrected GA underwent an O2 challenge test to define BPD.

Data Collection

HR and SpO2 were downloaded per pulse oximeter manufacturer instructions. Resuscitation interventions and physiologic responses were recorded at 30-second intervals by the OB circulating nurse per routine practice. Cases were reviewed continuously for adherence to NRP and study protocols.

Sample Collection and Analysis

One mL of umbilical cord blood and 0.25 mL of arterial blood were collected without anticoagulant on NICU admission. Serum was separated immediately by using centrifugation and stored at −80°C. All samples were analyzed within 1 month of collection. Samples were thawed carefully 1 at a time at room temperature, with mixing during and after the thawing process until equilibrium was reached. Thawed samples were analyzed immediately for TH and BAP with a Free Radical Analytical System (FRAS4 analyzer, H&D Srl, Parma, Italy) per manufacturer guidelines. TH represents the total of radical O2 metabolites produced by peroxidation of protein, lipids, and amino acids; it measures oxidative damage and serves as a biomarker of overall free radical attack. BAP measures both endogenous and exogenous antioxidative capacity of serum to reduce oxides by inactivating and eliminating free radicals and reactive O2 species. Although cytoplasmic antioxidants are not measured, BAP, by exploiting the chemical principle of the well-known ferrocyanide-reducing ability of plasma, provides a reliable measure of biological antioxidative potential of blood plasma. To estimate global balance between oxidative stress and antioxidant potential, the oxidative balance ratio (BAP/TH) was calculated based on measured values of BAP and TH.

Outcomes

The primary hypothesis was that LOX would improve oxidative balance ratio by 15% compared with HOX. Secondary outcomes were defined a priori as: reduction in admission PaO2 >100 mm Hg, change in Apgar score, total O2 use during resuscitation, integrated excessive O2 (Σ[FIO2 – 0.21] × time [min]), time during resuscitation with saturations >94%, use of rescue high-frequency oscillatory ventilation (HFOV), ventilator and hospital days, in-hospital mortality, and incidence of BPD, ROP, necrotizing enterocolitis, IVH, and PVL.

Statistical Analysis

Statistical analyses were conducted by using SAS version 9.2 (SAS Institute, Cary, NC). Normally distributed variables were compared by using Student’s t test. Nonparametric variables were compared by using Mann-Whitney rank sum tests. HR, SpO2, and FIO2 were compared with a linear mixed model repeated measures analysis, which included factors to assess group, time, and group-by-time interaction. Based on the reported oxidative balance ratio data for preterm neonates, to detect a 15% change, 39 patients were needed for each arm using a 2-sided α level of 0.05 and a power of 0.8.10 To accommodate a 10% potential patient dropout, 44 neonates were enrolled in each group.

RESULTS

Eligible neonates, comprising those enrolled and randomly assigned to LOX (n = 44) and HOX (n = 44), are shown in Fig 1. Fifteen neonates <28 weeks’ GA and 29 neonates 29 to 34 weeks’ GA were included in each arm. Eleven pairs of twins were included. One pair of monochorionic diamniotic twins were assigned randomly to HOX and 2 pairs of monochorionic diamniotic twins were assigned to LOX. For the remaining 8 pairs, each twin was randomly assigned to different groups. In 1 instance, only 1 twin required resuscitation and was enrolled. LOX and HOX neonates had similar maternal and baseline neonatal characteristics except that breech presentation was higher for HOX neonates (Table 2).

DR Resuscitation Interventions

As determined by the study protocol, the initial FIO2 was different for LOX and HOX neonates (Table 3). Use and maximum level of CPAP, need for PPV, maximum peak inspiratory pressure, and rates of intubation were similar. No neonate required chest compression.
or epinephrine. There was no difference in arterial cord pH, base deficit, or Apgar scores at 1 and 5 minutes. Two LOX infants and 3 HOX infants had an HR <60 beats per minute after 90 seconds of resuscitation, all of whom responded to ventilation via endotracheal tube. HR did not differ between groups at any time during resuscitation.

**DR O₂ Use**

LOX neonates were exposed to less O₂ in the DR as noted by lower O₂ load and integrated excessive O₂ compared with HOX (Table 3). Median FiO₂ and interquartile range for the LOX group was increased in a stepwise manner to 0.50 (0.30–0.70) by 6 minutes of life and subsequently weaned to 0.30 (0.21–0.45) by 10 minutes (Fig 2). Conversely, FiO₂ for HOX neonates was reduced in a stepwise manner to 0.70 (0.30–1) at 7 minutes of life and then further reduced to 0.35 (0.21–0.70) at 10 minutes. FiO₂ differed between LOX and HOX for the first 9 minutes of life. All LOX neonates required some increase in FiO₂ above room air to meet target SpO₂. Initial SpO₂ achieved during the first 2 minutes of life did not differ between groups. HOX neonates had higher SpO₂ between 3 and 6 minutes of life, but after 6 minutes, the results were similar between groups. More HOX neonates had SpO₂ >94% at each minute of life for the first 10 minutes compared with LOX (Fig 3). More LOX infants had saturations below the 10th centile for the first 10 minutes of life compared with HOX infants.²⁸

**Respiratory Support and Other Characteristics on Admission to NICU**

Respiratory support, including use and level of CPAP, mechanical ventilation, and peak inspiratory pressure were similar between LOX and HOX infants on NICU admission. Both groups had similar FiO₂, PaO₂, and SpO₂. Similar numbers of neonates had PaO₂ >100 mm Hg according to admission arterial blood gas levels.

**Oxidative Stress Markers and Antioxidant Capacity**

Cord blood TH (10⁷ [29–138] vs 10⁸ [85–161] U.CARR; P = .05), BAP (1649 [1415–1973] vs 1572 [1460–1776] mM; P = .05), and oxidative balance ratio (14 [11–19] vs 15 [9–18] mM/U.CARR; P < .05) were similar for LOX versus HOX. Within 1 hour of admission, TH was lower (10⁷ [91–146] U.CARR vs 151 [124–210] U.CARR; P = .001), BAP was higher (1564 [1216–1963] mM vs 1297 [1029–1705] mM; P = .02), and the oxidative balance ratio (13 [9–16] mM/U.CARR vs 8 [6–9] mM/U.CARR; P < .001) was higher for LOX versus HOX infants. When change from baseline (Δ) was analyzed, ΔTH was lower and Δoxidative balance ratio was higher for LOX infants, with no difference in ΔBAP (Fig 4).

**Respiratory and Other Short-Term NICU Morbidities**

Respiratory and other short-term morbidities are shown in Table 4. There were no differences in the incidence of...
respiratory distress syndrome, need for surfactant, pneumothorax, pulmonary interstitial emphysema, pulmonary arterial hypertension, pneumomediastinum, or pulmonary hemorrhage.

BPD according to the National Institutes of Health consensus and physiologic definition was lower for LOX infants, with a number needed to treat of 5 (95% confidence interval: 3–37) to prevent 1 case of BPD. LOX neonates received less rescue HFOV and spent fewer days on mechanical ventilation than HOX neonates. There were no differences in corticosteroid use for BPD, days on O2, maximum FiO2 during NICU stay, culture-proven sepsis, IVH grade 3 or 4, PVL, necrotizing enterocolitis, symptomatic patent ductus arteriosus, ROP stage 3 to 4, need for laser therapy, length of NICU stay, or mortality.

**DISCUSSION**

We report the first trial comparing LOX using goal saturations now recommended by the 2010 NRP guidelines versus HOX to demonstrate feasibility and decrease O2 exposure in preterm neonates.42 LOX successfully resuscitated preterm neonates without increasing the need for additional respiratory or cardiovascular support in the DR and decreased O2 load by one-half. LOX neonates spent less time with DR SO2 >94%, developed less oxidative stress by 1 hour of life, and had better preservation of BAP. Importantly, LOX also decreased ventilator days, rescue HFOV, and BPD.

Similar to previous small trials comparing various O2 strategies for DR resuscitation, the current study suggests that static delivery of 21% O2 is unlikely to achieve current NRP goal saturations for preterm infants. The majority needed supplemental O2 to meet these goals. Two previous DR O2 studies concluded that initiation of resuscitation with 21% O2 was inappropriate because the majority of preterm infants needed O2 supplementation to maintain the goal saturations prescribed by their study protocol. Our findings challenge this conclusion. Although it should not be expected that static delivery of room air will be adequate throughout resuscitation and stabilization, starting with 21% O2 provided it is gradually increased by using LOX, results in successful stabilization with no increase in sustained bradycardia, need for PPV, intubation, cardiovascular compressions, or vasoactive drugs.

NRP’s recommendation for the inter-quartile range of preductual saturations of healthy term infants as target goal saturations was based on consensus expert opinion as the United States took its first steps to stop the routine use of 100% O2 during resuscitation. The optimal SpO2 percentile for goal saturations during preterm transition is unknown, but ≥10th percentile is considered normal for parameters such as birth weight. Although in the first 10 minutes of life, the majority of LOX neonates had an SpO2 above the 10th percentile on the Dawson curves, more LOX infants had an SpO2 below the 10th percentile compared with HOX.42 Goos et al recently reported retrospectively that preterm neonates initially resuscitated with 30% O2 and adjusted to meet NRP goal saturations also resulted in some with SpO2 below the 10th percentile of the Dawson curves. The significance of this observation is speculative because the effect of brief DR LOX on long-term

**TABLE 3** DR Interventions and Early Parameters on Admission to NICU

<table>
<thead>
<tr>
<th>Parameter</th>
<th>LOX (n = 44)</th>
<th>HOX (n = 44)</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Initial FiO2</td>
<td>0.21 ± 0.08</td>
<td>1.0 ± 0.10</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>CPAP only, n (%)</td>
<td>11 (25)</td>
<td>11 (25)</td>
<td>NS</td>
</tr>
<tr>
<td>Maximum CPAP, cm H2O</td>
<td>5 (5–7)</td>
<td>5 (5–8)</td>
<td>NS</td>
</tr>
<tr>
<td>PPV, n (%)</td>
<td>29 (66)</td>
<td>30 (68)</td>
<td>NS</td>
</tr>
<tr>
<td>Maximum peak inspiratory pressure, cm H2O</td>
<td>25 (20–25)</td>
<td>25 (20–30)</td>
<td>NS</td>
</tr>
<tr>
<td>Intubation, n (%)</td>
<td>9 (20)</td>
<td>17 (39)</td>
<td>NS</td>
</tr>
<tr>
<td>Chest compression, n (%)</td>
<td>0</td>
<td>0</td>
<td>NS</td>
</tr>
<tr>
<td>Epinephrine, n (%)</td>
<td>0</td>
<td>0</td>
<td>NS</td>
</tr>
<tr>
<td>Arterial cord pH</td>
<td>7.25 ± 0.08</td>
<td>7.23 ± 0.10</td>
<td>NS</td>
</tr>
<tr>
<td>Arterial cord base deficit</td>
<td>6 ± 3</td>
<td>7 ± 5</td>
<td>NS</td>
</tr>
</tbody>
</table>

NS, not significant.

* Mean ± SD.

* Median (range).
neurodevelopmental outcome remains unknown. Studies comparing different starting concentrations of O2 to achieve target goal saturations and, more importantly, effects on both short- and long-term clinical outcomes are warranted; however, based on published studies so far, it seems unlikely that 21% vs 30% O2 would result in significant differences.

To optimize balance between excessive O2 and hypoxia, future studies comparing different target SpO2 ranges should be considered. We speculate that various strategies which improve ventilation and establishes functional residual capacity early may further reduce the need for supplemental oxygen.

The decrease in ventilator days, rescue HFOV, and BPD in LOX neonates is interesting; however, our study was not powered for secondary outcomes. Vento et al also reported decreased BPD with the use of a different LOX protocol. Although multiple factors promote lung injury resulting in BPD, oxidant injury is an important contributor. High O2 exposure can arrest lung septation in the saccular stage of development. Infants with BPD exposed to higher supplemental O2 to achieve higher levels of SpO2 have more persistent lung disease. We speculate that excess O2 exposure in the first few minutes of life may blunt vascular development in preterm infants. Development of strategies to minimize oxygen exposure to prevent tissue injury must be a priority.

![Figure 2](image-url)

**FIGURE 2**
A, Median level of O2 administered per minute of life for the first 10 minutes of life in preterm neonates randomly assigned to the LOX group, in which initial FiO2 after birth was 21%, or the HOX group, in which initial FiO2 after birth was 100%. *P < .05. B, Median SpO2 per minute of life for first 10 minutes of life in the DR in LOX and HOX infants. *P < .05.

![Figure 3](image-url)

**FIGURE 3**
A, Proportion of neonates with SpO2 >94% per minute for the first 10 minutes of life in the DR in the LOX and HOX groups. *P < .05. B, Proportion of neonates with SpO2 <10th percentile of Dawson curves for the first 10 minutes of life in the DR in the LOX and HOX groups. *P < .05.
A recent meta-analysis comparing lower (21%–50%) versus higher (>50%) O₂ concentrations for DR resuscitation of preterm infants and their effects on morbidity and mortality identified 6 randomized trials. Lower O₂ reduced the risk of death, but this effect disappeared when limited to studies with adequate allocation concealment. The authors stressed the need for larger trials powered to detect important outcomes, including neurodevelopment and mortality.

The current study has several limitations. Although randomized, the trial was not blinded and is at-risk for bias because the resuscitation team was aware of treatment assignments. However, DR interventions were diligently recorded by the OB circulating nurse who was not involved in the trial (as is our standard practice), and systematic errors that might have unduly influenced randomization or outcome were not found. Video recordings or computerized data acquisition systems other than pulse oximetry for continuous physiologic data were not available. The number of neonates recruited to the trial, which was powered for a difference in oxidative stress markers, did not allow for the evaluation of short-term clinical outcomes such as BPD or long-term neurodevelopmental outcomes with sufficient statistical power.

One of the strengths of the study is that it recapitulates the intended real-world DR use of O₂. Adjusting FIO₂ in response to SpO₂ every 30 seconds is challenging and might interfere with or delay other steps needed to resuscitate a preterm infant. Lack of research personnel assigned to make these changes provided opportunity to study the effect of these complex interactions in a randomized controlled trial setting. We did not have a visual O₂ targeting system such as TOTS [Transitional Oxygen Targeting System] to help keep saturations within target range. Our study is relevant to most resuscitation situations because the majority of the world’s neonates are resuscitated without a visual targeting system. This is the first randomized trial to use current NRP-recommended O₂ saturation targets in preterm infants needing DR resuscitation. Unlike previous trials examining resuscitation of preterm neonates with LOX, our study included all eligible preterm infants due to antenatal waiver of consent, thereby reducing selection bias and increasing the generalizability of the results.

FIGURE 4
(A) Total hydroperoxides, (B) biological antioxidant potential, and (C) oxidative balance ratio in cord blood and 1 hour of life in preterm neonates resuscitated with LOX or HOX. Data are presented as mean ± SD. Mann-Whitney test was used for comparison of difference over time (cord and admission samples) between the groups. *P < .01.

### TABLE 4 Respiratory and Other Short-Term Morbidities

<table>
<thead>
<tr>
<th>Parameter</th>
<th>LOX (n = 44)</th>
<th>HOX (n = 44)</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Respiratory morbidities</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Respiratory distress syndrome</td>
<td>28 (64)</td>
<td>25 (57)</td>
<td>NS</td>
</tr>
<tr>
<td>Received surfactant</td>
<td>15 (34)</td>
<td>17 (39)</td>
<td>NS</td>
</tr>
<tr>
<td>Pneumothorax</td>
<td>2 (4)</td>
<td>2 (4)</td>
<td>NS</td>
</tr>
<tr>
<td>Pulmonary interstitial emphysema</td>
<td>2 (4)</td>
<td>5 (11)</td>
<td>NS</td>
</tr>
<tr>
<td>Pulmonary arterial hypertension</td>
<td>1 (2)</td>
<td>3 (7)</td>
<td>NS</td>
</tr>
<tr>
<td>Pneumomediastinum</td>
<td>2 (4)</td>
<td>0 (0)</td>
<td>NS</td>
</tr>
<tr>
<td>Pulmonary hemorrhage</td>
<td>2 (4)</td>
<td>6 (14)</td>
<td>NS</td>
</tr>
<tr>
<td>BPD</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>O₂ at 28 d</td>
<td>8 (19)</td>
<td>17 (39)</td>
<td>.04</td>
</tr>
<tr>
<td>O₂ at 36 wk</td>
<td>3 (7)</td>
<td>11 (25)</td>
<td>.04</td>
</tr>
<tr>
<td>Physiologic definition</td>
<td>2 (4)</td>
<td>10 (23)</td>
<td>.03</td>
</tr>
<tr>
<td>Intravenous steroids for BPD</td>
<td>1 (2)</td>
<td>2 (4)</td>
<td>NS</td>
</tr>
<tr>
<td>HFOV</td>
<td>2 (4)</td>
<td>10 (23)</td>
<td>.03</td>
</tr>
<tr>
<td>Days on mechanical ventilation</td>
<td>3 (0–46)</td>
<td>8 (0–96)</td>
<td>.045</td>
</tr>
<tr>
<td>Days on CPAP</td>
<td>4 (0–44)</td>
<td>6 (0–43)</td>
<td>NS</td>
</tr>
<tr>
<td>Days on oxygen</td>
<td>4 (0–113)</td>
<td>10 (0–163)</td>
<td>NS</td>
</tr>
<tr>
<td>Maximum FO₂ b</td>
<td>64 (25–100)</td>
<td>100 (26–100)</td>
<td>NS</td>
</tr>
<tr>
<td>Other short-term morbidities</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sepsis proven</td>
<td>8 (18)</td>
<td>10 (23)</td>
<td>NS</td>
</tr>
<tr>
<td>IVH grade 3 or 4</td>
<td>1 (2)</td>
<td>1 (2)</td>
<td>NS</td>
</tr>
<tr>
<td>Cystic PVL</td>
<td>0 (0)</td>
<td>1 (2)</td>
<td>NS</td>
</tr>
<tr>
<td>Necrotizing enterocolitis</td>
<td>1 (2)</td>
<td>6 (14)</td>
<td>NS</td>
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<tr>
<td>Symptomatic patent ductus arteriosus</td>
<td>6 (14)</td>
<td>10 (23)</td>
<td>NS</td>
</tr>
<tr>
<td>Severe ROP/need for laser therapy</td>
<td>1 (2)</td>
<td>4 (9)</td>
<td>NS</td>
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<tr>
<td>Days of hospitalization</td>
<td>33 (7–126)</td>
<td>40 (8–166)</td>
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</tr>
<tr>
<td>Death during initial hospitalization</td>
<td>2 (4)</td>
<td>3 (7)</td>
<td>NS</td>
</tr>
</tbody>
</table>

NS, not significant.

* The physiologic definition of BPD includes, as a criterion, the receipt of >30% O₂ or the need for positive pressure support at 36 weeks or, in the case of infants requiring <30% O₂, the need for any O₂ at 36 weeks after an attempt at O₂ withdrawal.

* Median (range).
28-week and 29- to 34-week GA. This approach is important because of limited data on the impact of LOX on more mature preterm neonates.

CONCLUSIONS
LOX for DR resuscitation of preterm neonates is feasible, decreases O₂ exposure without increasing need for additional resuscitation, and decreases oxidative stress. In addition, LOX seems to decrease ventilator days, HFOV, and risk of BPD. Thus, LOX is a reasonable alternative for DR resuscitation of preterm neonates. Large randomized trials adequately powered for evaluation of the relationship between O₂ load during resuscitation and important clinical outcomes such as BPD and long-term neurodevelopment are warranted.53

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Resuscitation of Preterm Neonates With Limited Versus High Oxygen Strategy
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