Bi-level CPAP does not improve gas exchange when compared with conventional CPAP for the treatment of neonates recovering from respiratory distress syndrome

Andrea L Lampland,1,2 Brenda Plumm,1 Cathy Worwa,1 Patricia Meyers,1 Mark C Mammel1,2

ABSTRACT

Aim We hypothesised that short-term application of bi-level nasal continuous positive airway pressure CPAP (SiPAP) compared with conventional nasal CPAP (nCPAP) at the same mean airway pressure in infants with persistent oxygen need recovering from respiratory distress syndrome would improve CO2 removal with no change in oxygen requirement.

Design Non-blinded, randomised, observational four-period crossover study.

Setting/population Level III NICU; low-birthweight infants requiring CPAP and oxygen while recovering from respiratory distress syndrome.

Methods Infants requiring nasal CPAP for >24 h prior to study enrolment, and fraction of inspired oxygen requirement (FiO2) of 0.25–0.5, were randomised to either nCPAP or SiPAP. A crossover design with four 1 h treatment periods was used such that each infant received both treatments twice. Oxygen saturations (SaO2), transcutaneous CO2 (tcCO2) and vital signs were monitored continuously. Polysomnographic recordings were analysed for apnoea, bradycardia and oxygen desaturation.

Results Twenty low-birthweight infants receiving 0.3 ±0.04% supplemental oxygen on CPAP of 6 cm H2O were studied at an average of 33 days of age (±23 days, SD). There were no differences in tcCO2 or other physiological parameters except mean blood pressure, which was lower during nCPAP (52.3±8.3 vs 54.4±9.1 mm Hg; ±SD; p<0.01). No differences in short or prolonged apnoea, bradycardia or significant desaturation events were observed.

Conclusions At similar mean airway pressures, SiPAP does not improve CO2 removal, oxygenation or other studied physiological parameters with the exception of mean blood pressure, which was not clinically significant.

Trial registration number NCT01053455.

INTRODUCTION

Respiratory distress syndrome (RDS) remains a leading cause of morbidity and mortality in premature infants.1 The cascade of events that typifies RDS and its long-term counterpart, chronic lung disease, is rooted in the intrinsic deficits of the premature lung as well as exacerbated by mechanical ventilation.2 3 Non-invasive ventilatory strategies, such as nasal continuous positive airway pressure (nCPAP), minimise lung inflammation and injury associated with mechanical ventilation.4 Avoidance of intubation and increased use of early nCPAP to treat RDS has been shown to decrease exposure to mechanical ventilation and decrease the duration of supplemental oxygen therapy.5–8 In babies that require intubation and mechanical ventilation, use of nCPAP at the time of extubation has also been shown to decrease extubation failure.9

Recent studies have documented that the use of non-invasive nasal intermittent mechanical ventilation (NIPPV), with short inspiratory times and adequate ventilation rates, can satisfactorily treat hypventilation, apnoea of prematurity and potentially decrease extubation failure when compared with the use of nCPAP.10 11 However, most published studies of NIPPV in neonates use a very broad definition to include any form of nCPAP that
has an intermittent increase in applied pressure. One type of NIPPV is bi-level CPAP where CPAP delivery systems cycle the positive airway pressure between two levels and allow patients to breathe throughout the respiratory cycle, with the potential to improve oxygenation and ventilation. However, there have been very few studies to date looking at the use of bi-level CPAP as compared with the standard use of nCPAP in premature infants with respiratory distress, and those that exist have differences in how the bi-level CPAP support is delivered.12–14 To gain clarity on this commonly used mode of NIPPV, we compared unsynchronised bi-level nasal CPAP (Infant Flow SiPAP System, CareFusion Corporation, San Diego, California, USA) with standard nasal CPAP in low-birthweight infants with respiratory distress.

METHODS
In this randomised, observational crossover study, patient eligibility was based on the following inclusion criteria: birth weight <2500 g, ongoing treatment of respiratory distress with nasal CPAP for >24 h prior to study enrolment and fraction of inspired oxygen requirement (FiO2) of 0.25–0.5 to keep oxygen saturations 85–95% for a minimum of 1 h prior to initiation of the study. Exclusion criteria included FiO2 requirement of <0.25 or >0.5; active medical treatment for PDA or culture proven sepsis, congenital defects of the airway, lungs or oesophagus, congenital cyanotic heart defects, genetic syndromes, or postoperative recovery period of <24 h. This study was registered at http://www.clinicaltrials.gov (#NCT01053455) and approved by the Children’s Hospitals and Clinics of Minnesota Institutional Review Board.

After written informed consent was obtained, the patient was placed in the supine position with the mouth closed by the aid of a soft chinstrap per our NICU standard care. A transcutaneous carbon dioxide monitor as well as pneumocardiogram sensors were placed on the infant (SenTec Digital Monitoring System, Therwil, Switzerland; Philips SmartMonitor, Andover, Massachusetts, USA; SpaceLabs Inc, Redmond, Washington, USA). The patient was randomised by sealed envelope shuffle to a starting treatment mode of either nCPAP at 6 cm H2O pressure (6 cm H2O) with a minimum difference in upper and lower pressure levels set to deliver the same mean airway pressure or unsynchronised bi-level CPAP (SiPAP) with the upper and lower pressure levels set to deliver the same mean airway pressure (6 cm H2O) with a minimum difference in upper and lower pressures of 3 cm H2O. SiPAP was set with the upper pressure to be delivered for 1 s at a rate of 20 cycles to the upper pressure per minute. Hudson short binalar prongs (Hudson Respiratory Care, Temecula, California, USA) were always used. All support was delivered by the Infant Flow SiPAP System with a flow rate of 8–9 L/min in either the CPAP or BiPhasic mode (CareFusion Corporation, San Diego, California, USA). Research personnel adjusted the FiO2 to attain a targeted oxygen saturation of 88–90%. The patients were maintained in the usual thermoneutral environment throughout the study, and all prescribed therapies were performed as ordered by the primary care team.

At study initiation, the infant was started on the randomised starting mode of either nCPAP or SiPAP. The study consisted of four 1 h study blocks, alternating from the initial mode to the alternate mode twice. During each study block, transcutaneous CO2 levels (tcCO2), heart rate, respiratory rate and oxygen saturations were measured continuously (data acquisition every second using SenTec Digital Monitoring System, Therwil, Switzerland; SpaceLabs Inc, Redmond, Washington, USA). Three-channel pneumocardiogram data, including heart rate, respiratory effort, SaO2 and chest wall impedance, were collected continuously for each block along with a nursing event log of patient cares/events (data acquisition every second using Philips SmartMonitor, Andover, MA). Apnoeic episodes were defined as absence of thoracic impedance change for a minimum of 10 s. Bradycardic episodes were defined as persistent heart rate <80 beats per minute for a minimum of 10 s. Significant desaturation episodes were defined as persistent pulse oximetry values <80% for a minimum of 10 s. Manual blood pressures were taken with appropriate sized neonatal blood pressure cuff every 10 min throughout the study. The study ended when the patient completed the 4 h study or was terminated early if the patient developed any signs of intolerance during the study, including persistent tachypnoea (respiratory rate >80 breaths per minute for >10 consecutive minutes), an increase of >50% in the number of episodes of apnoea or bradycardia compared with the prestudy baseline noted 1 h preceding study entry, or increased supplemental FiO2 >0.3 from prestudy baseline.

To allow for equilibration, we grouped and analysed data points from the last 20 min of each treatment block. Pneumocardiogram data were independently analysed and scored by a polysomnographic technologist blinded to study group. A sample size of 17 was calculated to detect a mean difference of 3 mmHg tcCO2 based on a two-tailed p value of 0.05, power of 0.9 and a within-patient SD of 2.5 mm Hg. Data were analysed using commercial statistical software (Graphpad Prism V3.0a; Chicago, Illinois, USA). Paired two-tailed t tests were employed, and p values <0.05 were considered statistically significant.

RESULTS
Twenty low-birthweight (<2500 g) infants were enrolled (table 1). Eleven mothers received antenatal betamethasone treatment prior to delivery. In total, 17 of the 20 infants were delivered by caesarean section. Also, 19 of the 20 infants had a history of previous endotracheal intubation, exogenous surfactant administration and mechanical ventilation prior to study enrolment. All infants received caffeine citrate therapy for treatment of apnoea of prematurity. The average supplemental oxygen requirement at time of randomisation was FiO2 of 0.3 (±0.04). During the SiPAP blocks, the setting for the average high pressure was 8.9 cm H2O (±0.4 cm H2O, SD) and the average low pressure was 4.1 cm H2O (±0.2 cm H2O).

There were no differences in tcCO2 values among the two treatment groups. Physiological variables, including heart rate, oxygen saturation, supplemental oxygen needs and respiratory rate, were similar among treatment groups. Average diastolic and mean blood pressure values were significantly higher in the SiPAP group compared with the nCPAP group (table 2). There

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Patient demographics</th>
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<tbody>
<tr>
<td>Mean value</td>
<td>SD</td>
</tr>
<tr>
<td><strong>Gestational age</strong></td>
<td>26 weeks</td>
</tr>
<tr>
<td></td>
<td>5 days</td>
</tr>
<tr>
<td><strong>Days of life at study entry</strong></td>
<td>33 days</td>
</tr>
<tr>
<td><strong>Birth weight</strong></td>
<td>897 g</td>
</tr>
<tr>
<td><strong>Weight at study entry</strong></td>
<td>1310 g</td>
</tr>
<tr>
<td><strong>Days on nCPAP before study entry</strong></td>
<td>7.6 days</td>
</tr>
</tbody>
</table>

For all infants, the date of birth was considered day of life 1. nCPAP, nasal continuous positive airway pressures.
were no significant differences in short (10–19 s) or prolonged (≥20 s) apnoeic, bradycardic or desaturation events between the two treatment groups (table 3).

**DISCUSSION**

In this randomised crossover trial, we compared two strategies for non-invasive support in babies recovering from RDS. Nasal CPAP is a strategy intended to recruit and stabilise lung volumes, and by doing so improve the mechanical behaviour of the lung. Bi-level CPAP is a newer non-invasive respiratory support technique that is essentially CPAP delivered at different pressures in a time-cycled manner, with ordered settings that are termed similarly to those used during invasive mechanical ventilation. In bi-level CPAP, the upper pressure settings are much lower than typical peak inspiratory pressures used with invasive mechanical ventilation and the bi-level CPAP inspiratory times are often much longer than those typically used with mechanical ventilation. This short-term crossover study was designed to test a simple physiological hypothesis that bi-level CPAP administered at the same mean airway pressure as nCPAP would improve ventilation and lower CO₂, as reflected in tcCO₂ values due to the cyclic pressure changes intended to augment minute ventilation.

In this study, we saw no evidence of improved ventilation during bi-level CPAP as the tcCO₂ values were not different between the two treatment groups. Choosing a sample size adequate to detect a 3 mm Hg change in tcCO₂ with 90% power ensured that any meaningful change, if present, could have been detected. We also found no evidence that bi-level CPAP significantly impacted oxygenation, apnoea, bradycardia or desaturation events. We did observe a statistically significant, but clinically unimportant, change in blood pressures, with slightly higher mean and diastolic blood pressure values recorded during the bi-level CPAP treatment periods. We speculate that the mean and diastolic blood pressures were higher in the bi-level CPAP group due to the 10 min sampling interval during the study blocks. Potentially, transient positive end-expiratory pressure-induced changes in cardiac output and blood pressure could have resulted in these differences by chance.¹³

The use of bi-level CPAP has gained wide acceptance in spite of limited information on its indications, efficacy and proper application.¹²–¹⁴ Three studies evaluating the use of bi-level CPAP in preterm infants have been published. All three studies describe their choice of settings for bi-level CPAP as empiric since optimal parameters of the bi-level pressures have not been investigated. Migliori and colleagues’ non-blinded crossover study in 2005 is most similarly structured to our study. In contrast to our study, Migliori’s study provided increased mean airway pressure during the bi-level CPAP treatment blocks and cycled to the upper-level pressure more frequently (30 times per minute for 0.5 s per cycle). They evaluated bi-level CPAP delivered by the Infant Flow Advance ventilator (Electro Medical Equipment, Ltd., Brighton, UK) to nasal CPAP in 20 low-birthweight infants and found a significant improvement in gas exchange during the bi-level CPAP treatment periods. They hypothesise that their findings of improved oxygenation and ventilation are related to increased mean airway pressure and ventilator-induced increase in tidal volume during the bi-level CPAP blocks. However, it still begs the question of whether the biphasic nature of the bi-level CPAP produces these effects or whether application of nasal CPAP at higher mean airway pressure would result in similar improvements.

Two other more recent studies have attempted to evaluate short-term safety and efficacy of bi-level CPAP compared with standard nasal CPAP. In 2010, Lista and colleagues compared synchronised bi-level CPAP and standard nasal CPAP in infants 28–34 weeks’ gestation with respiratory distress at 1 h of life. They found no differences in short-term markers of inflammation between the two treatments. Although not powered for long-term pulmonary endpoints, their study did find the infants treated with bi-level CPAP had a significantly shorter duration of respiratory support and supplemental oxygen need than the group randomised to standard CPAP. In 2012, O’Brien and colleagues performed a randomised controlled trial of bi-level CPAP versus standard nasal CPAP to facilitate sustained extubation in babies ≤1250 g. Although the trial was stopped five infants short of reaching the calculated sample size, they found that the use of bi-level CPAP was as effective as, but not statistically better than, standard CPAP at aiding sustained extubation.

Previous meta-analysis data have concluded that nasal intermittent positive pressure ventilation (NIPPV), of which bi-level CPAP is a subtype, increases the effectiveness of nCPAP by preventing the need for reintubation.¹⁰ However, this analysis is based on only three trials, performed between 1999 and 2002, using synchronised forms of NIPPV None used SiPAP; additionally, synchronised SiPAP is not approved for use in the USA. These studies do not address how NIPPV may impact gas exchange and produce its effects. A recent large multicentre randomised controlled trial compared nCPAP with NIPPV in 1009 infants. The study was designed to compare rates of death before 36 weeks adjusted age or survival with bronchopulmonary dysplasia, as well as a number of secondary outcomes. Contrary to the meta-analysis conclusions, this study found no evidence that NIPPV in extremely low-birthweight infants improved the primary outcome or any of the many secondary outcomes studied.¹⁷ The questions asked in this study were

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**Table 2** Primary physiological and respiratory variables

<table>
<thead>
<tr>
<th>Variable</th>
<th>CPAP</th>
<th>SiPAP</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Heart rate (bpm)</td>
<td>168.2 (±12.3)</td>
<td>167.3 (±12.1)</td>
<td>0.4</td>
</tr>
<tr>
<td>O₂ saturation (%)</td>
<td>87.4 (±2.9)</td>
<td>87.3 (±3.8)</td>
<td>1.0</td>
</tr>
<tr>
<td>Transcutaneous CO₂ (mm Hg)</td>
<td>54 (±7.1)</td>
<td>53.2 (±6.4)</td>
<td>0.9</td>
</tr>
<tr>
<td>Respiratory rate (bpm)</td>
<td>47.9 (±9.5)</td>
<td>47.6 (±9.9)</td>
<td>0.5</td>
</tr>
<tr>
<td>FiO₂ (%)</td>
<td>30.1 (±4.8)</td>
<td>29.4 (±4.3)</td>
<td>1.0</td>
</tr>
<tr>
<td>Systolic blood pressure (mm Hg)</td>
<td>68.4 (±10.4)</td>
<td>69.8 (±10.9)</td>
<td>0.1</td>
</tr>
<tr>
<td>Diastolic blood pressure (mm Hg)</td>
<td>42.5 (±7.4)</td>
<td>45.1 (±8.8)</td>
<td>0.003**</td>
</tr>
<tr>
<td>Mean blood pressure (mm Hg)</td>
<td>52.3 (±8.3)</td>
<td>54.4 (±9.1)</td>
<td>0.01**</td>
</tr>
</tbody>
</table>

Means±SD. **p value <0.05. bpm, beats per minute; brpm, breaths per minute; CPAP, continuous positive airways pressure.

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**Table 3** Pneumogram analysis

<table>
<thead>
<tr>
<th>Event</th>
<th>CPAP</th>
<th>SiPAP</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Apnoea 10–19 s</td>
<td>1.1 (±0.7)</td>
<td>1.2 (±0.1)</td>
<td>0.9</td>
</tr>
<tr>
<td>Bradycardia 10–19 s</td>
<td>0 (±0.2)</td>
<td>0.1 (±0.3)</td>
<td>0.3</td>
</tr>
<tr>
<td>Desaturation 10–19 s</td>
<td>2.0 (±2.3)</td>
<td>1.9 (±2.9)</td>
<td>1.0</td>
</tr>
<tr>
<td>Apnoea ≥20 s</td>
<td>0.4 (±1.1)</td>
<td>1.0 (±4.6)</td>
<td>0.5</td>
</tr>
<tr>
<td>Bradycardia ≥20 s</td>
<td>0.03 (±0.2)</td>
<td>0 (±0)</td>
<td>0.3</td>
</tr>
<tr>
<td>Desaturation ≥20 s</td>
<td>2.7 (±4.4)</td>
<td>2.9 (±5.0)</td>
<td>0.7</td>
</tr>
</tbody>
</table>

Mean number of episodes per study block ±SD. CPAP, continuous positive airways pressure.
global rather than targeted physiology outcomes and might seem to make the data we present moot. However, NIPPV in this pragmatic study was delivered using a number of different techniques, including bi-level CPAP as well as synchronised and unsynchronised ventilator-derived NIPPV. The question remains whether any of these techniques studied individually, as we have done, might produce different results, either acutely or over the long term.

Limitations of our study include its small sample size, heterogeneity in patient age and length of time on nCPAP before entering the study. Our population was recovering from RDS and requiring moderate supplemental oxygen on nCPAP, but overall, they were stable. Our analysis of only short-term physiological outcomes prevents any conclusions about important longer-term problems. Though only 20 patients were studied, the crossover design is powerful and allowed easy assessment of our targeted short-term respiratory outcomes. The addition of other important short-term respiratory physiology assessments, such as measurements of infant work of breathing in each treatment mode, would add further information. We acknowledge that short-term physiology does not define long-term outcomes but is an important first step for understanding how and why new techniques might be useful.

We studied unsynchronised SiPAP. Just as synchronisation produces important differences during convention mechanical ventilation, there may be differences between unsynchronised and synchronised non-invasive techniques. However, little information is available and is conflicting as to whether synchronisation confers any significant benefit. We chose unsynchronised SiPAP since it is the only type of bi-level CPAP available in the USA and is commonly used in other parts of the world. We acknowledge that by keeping mean airway pressure the same between periods, we used SiPAP in a manner that may differ from some physicians’ clinical practice; at least some physicians use SiPAP at higher mean airway pressures than that used during nCPAP. We chose to keep mean airway pressure the same to minimise positional changes in lung volumes and oxygenation that might result and to better identify if the variable nature of pressure exposure by SiPAP in and of itself allows for improved short-term respiratory outcomes.

In conclusion, we compared two non-invasive respiratory support techniques, nCPAP and SiPAP, in a randomised crossover study in babies requiring oxygen therapy and recovering from RDS. We hypothesised that, at similar mean airway pressures, SiPAP would lower tCO2. There were no differences in tCO2 values between the two treatment groups or in other studied variables except for blood pressure. We saw a small but significant change in mean and diastolic blood pressures during SiPAP. Using this technique, SiPAP provides no physiological benefit compared with nCPAP in acute gas exchange or respiratory stability.

Contributors All, BP, CW, PM and MCM made substantive contributions to designing and implementing this clinical study and agree to be accountable for all aspects of the work they have done. BP, CW and PM made substantial contributions to data acquisition and analysis, critically revising the manuscript and giving final approval to the manuscript. All and MCM made substantial contributions to data analysis and interpretation, drafting and critically revising the manuscript and approving the final manuscript.

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Competing interests AL is a clinical research steering committee member for Discovery Labs, Inc.

Ethics approval Children’s Hospitals and Clinics of Minnesota IRB.

Provenance and peer review Not commissioned; externally peer reviewed.

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