Initiating resuscitation before umbilical cord clamping in infants with congenital diaphragmatic hernia: a pilot feasibility trial

Elizabeth E Foglia, Anne Ades, Holly L Hedrick, Natalie Rintoul, David A Munson, Julie Moldenhauer, Juliana Gebb, Bonnie Serletti, Aasma Chaudhary, Danielle D Weinberg, Natalie Napolitano, María Victoria Fraga, Sarah J Ratcliffe

ABSTRACT

Background Infants with congenital diaphragmatic hernia (CDH) often experience hypoxaemia with acidosis immediately after birth. The traditional approach in the delivery room is immediate cord clamping followed by intubation. Initiating resuscitation prior to umbilical cord clamping (UCC) may support this transition.

Objectives To establish the safety and feasibility of intubation and ventilation prior to UCC for infants with CDH. To compare short-term outcomes between trial participants and matched controls treated with immediate cord clamping before intubation and ventilation.

Design Single-arm, single-site trial of infants with CDH and gestational age ≥36 weeks. Infants were placed on a trolley immediately after birth and underwent intubation and ventilation, with UCC performed after qualitative CO₂ detection. The primary feasibility endpoint was successful intubation prior to UCC. Prespecified safety and physiological outcomes were compared with historical controls matched for prognostic variables using standard bivariate tests.

Results Of 20 enrolled infants, all were placed on the trolley, and 17 (85%) infants were intubated before UCC. The first haemoglobin and mean blood pressure values, vasoactive medications, inhaled nitric oxide or extracorporeal membrane oxygenation were all significantly higher in trial participants than controls. There were no significant differences between groups for subsequent blood pressure values, vasoactive medications, inhaled nitric oxide or extracorporeal membrane oxygenation. Blood gas and oxygenation index values did not differ between groups at any point.

Conclusions Intubation and ventilation prior to UCC is safe and feasible among infants with CDH. The impact of this approach on clinically relevant outcomes deserves investigation in a randomised trial.

INTRODUCTION

Physiological adaptation after birth is a critical transition for infants with congenital diaphragmatic hernia (CDH). In utero, the fetus has fluid-filled lungs, and gas exchange occurs at the placenta. Immediately after birth, the infant must aerate the lungs, which triggers a physiological transition of reduced pulmonary vascular resistance, increased pulmonary blood flow and gas exchange in the lung. Infants with CDH struggle to independently achieve lung aeration due to pulmonary hypoplasia and space-occupying effects of herniated abdominal organs. Furthermore, they are at risk for decreased pulmonary blood flow due to pulmonary hypertension. In a preclinical model of CDH, physiological challenges during transition include lower lung compliance, more respiratory acidosis and poor cerebral oxygenation. Clinically, this manifests as hypoxaemia, hypercarbia and acidosis after birth, often reflected in low Apgar scores.

Limited data inform the delivery room management for infants with CDH. The standard approach is immediate umbilical cord clamping (UCC) followed by intubation and ventilation. An alternative strategy is to establish lung aeration prior to UCC, which has been called ‘physiologically based cord clamping’. Physiologically based cord clamping may stabilise gas exchange during neonatal transition among infants with CDH by supporting aeration of the hypoplastic lung and increasing pulmonary blood flow through the thickened pulmonary vasculature before UCC. In
METHODS

Study design and setting

This was a single-arm, single-site trial at the Children’s Hospital of Philadelphia (ClinicalTrials.gov identifier: NCT03314233). Eligible infants had an antenatal diagnosis of CDH and were at least 36 weeks’ gestation at birth. Exclusion criteria were: other major anomalies, enrolment in an ongoing trial of fetoscopic endoluminal tracheal occlusion, multiple gestation, palliative care planned or considered, placental abnormalities (abruption, previa or accreta) and maternal magnesium sulfate therapy. Informed consent was obtained from mothers of potentially eligible infants in the prenatal clinic setting, and eligibility criteria were confirmed immediately before delivery. If all eligibility criteria were met, the mother/infant dyad was allocated to receive the study intervention.

Intervention

An attending neonatologist, respiratory therapist and neonatal nurse attended each delivery (Figure 1). A mobile LifeStart Trolley (Inditherm Medical, Rotherham, UK) was adjusted to the level of the introitus for vaginal deliveries and just above the level of the incision for caesarean deliveries. An activated chemical warmer mattress was placed on the trolley mattress for thermoregulation. For caesarean deliveries, the trolley and neonatal providers were sterilely draped.

Initially, the study used a purpose-built respiratory pole with air and oxygen tanks and a Neopuff Infant T-piece resuscitator (Fisher and Paykel, Auckland, New Zealand). Suction tubing was connected to canisters in the labour room or obstetrical suite. Midway through the trial, this equipment was replaced by a portable T-piece resuscitator that included suction (Giraffe Stand Alone Resuscitation System, GE Healthcare, USA). Airway equipment and respiratory tubing were not sterile and were handled to avoid contact with the sterile operative field for caesarean deliveries.

Immediately after delivery, the obstetrical provider placed the infant on the trolley. The neonatologist intubated the infant, and ventilation was commenced using settings per hospital protocol (initial pressures 20–25/5 cm H₂O, fractional inspired oxygen 0.5). UCC was performed after consistent qualitative colorimetric CO₂ detection, with guidelines for maternal uterotonetic administration after UCC. After UCC, the neonatal team moved the trolley and respiratory equipment away from the mother, taped the endotracheal tube, covered the infant with warm towels and transported the infant with ongoing ventilation to the resuscitation warmer bed. All remaining care was per clinical guidelines. The protocol stipulated for UCC to be performed prior to intubation for any of the following: the infant could not be placed on the trolley, intubation was not successful within 3 min from birth, cord avulsion or bleeding or any obstetrical or neonatal provider concerns.

The study team developed training videos and conducted equipment and simulation training for staff. A study team member was present for all resuscitations to ensure protocol adherence. When time allowed, the intervention was video recorded.

Outcomes

The primary feasibility endpoint was successful intubation prior to UCC within the 3 min timeframe. Prespecified neonatal safety outcomes included cord avulsion, cardiopulmonary resuscitation and first measured temperature. Maternal safety outcomes included estimated blood loss, therapeutic uterotonics and...
Physiological outcomes included first haemoglobin measurement, mean blood pressure values and blood gas parameters in the first 48 hours after birth, the first clinically obtained echocardiogram and mortality in the first week after birth. Echocardiogram images were independently reviewed by an unblinded single assessor (MVF) for signs of pulmonary hypertension. If present, the severity of pulmonary hypertension was graded based on the most severe assessment of the following criteria: direction of shunting at the ductus arteriosus (bidirectional: moderate; all right to left: severe), estimated systolic right ventricular pressures through continuous wave interrogation of the tricuspid regurgitant jet and calculation of a systolic right ventricle-to-right atrium pressure gradient by using the modified Bernoulli equation [pressure gradient=4× jet velocity²] (≤½ systemic: normal/mild; ≥½ and < systemic: moderate; ≥systemic: severe) and qualitative evaluation of interventricular septal position at the end of systole (rounded: normal/mild; flattened: moderate and bowed: severe).

Interventions assessed included respiratory support and inhaled nitric oxide in the first 48 hours of life and extracorporeal membrane oxygenation (ECMO) in the first week after birth. Historical controls were identified from a ongoing pulmonary hypoplasia programme registry and matched with trial participants on the basis of gestational age, mode of delivery, side of CDH, liver position (intrathoracic vs intra-abdominal) and observed to expected lung-to-head ratio (O/E LHR) obtained, and these images were not assessed. All infants had any portion of liver in chest, n (%) 14 (70) 13 (68) left sided defect, n (%) 15 (75) 15 (79) any portion of liver in chest, n (%) 14 (70) 13 (68) O/E LHR % median (IQR) 30.1 (25.6–35.9) 34.2 (27.4–36.7) *One trial participant not matched due to postnatal diagnosis of major anomaly. O/E LHR %, observed to expected lung-to-head ratio from anterior-posterior diameter method by the TOTAL trial calculator for infants with left-sided congenital diaphragmatic hernia.

Table 1 Baseline characteristics of trial participants

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Trial participants (n=20)</th>
<th>Historical controls (n=19)*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gestational age, weeks; mean (SD)</td>
<td>38.7 (0.5)</td>
<td>38.5 (0.8)</td>
</tr>
<tr>
<td>Birth weight, grams; mean (SD)</td>
<td>3374 (499)</td>
<td>3409 (577)</td>
</tr>
<tr>
<td>Male sex, n (%)</td>
<td>15 (75)</td>
<td>12 (63)</td>
</tr>
<tr>
<td>Vaginal delivery, n (%)</td>
<td>8 (40)</td>
<td>8 (42)</td>
</tr>
<tr>
<td>Left sided defect, n (%)</td>
<td>15 (75)</td>
<td>15 (79)</td>
</tr>
<tr>
<td>Any portion of liver in chest, n (%)</td>
<td>14 (70)</td>
<td>13 (68)</td>
</tr>
<tr>
<td>O/E LHR %; median (IQR)</td>
<td>30.1 (25.6–35.9)</td>
<td>34.2 (27.4–36.7)</td>
</tr>
</tbody>
</table>

RESULTS

Between January and October 2018, 20 eligible infants were allocated to receive the intervention (figure 2). Demographic characteristics of enrolled infants and matched controls are shown in table 1. One enrolled infant was diagnosed with a second major anomaly postnatally. This infant was included in the assessment of safety and feasibility but not physiological outcomes; no control was matched to this participant.

All 20 infants were placed on the trolley and 17 (85%) infants were successfully intubated prior to UCC (online supplementary video). The median interval between birth and UCC was 2 min (IQR 1:15–2:32). Among the 17 successful cases, the median duration between birth and intubation was 1:02 (IQR 0:55–2:00), with 14 infants intubated on the first attempt and 3 intubated on the second attempt within this timeframe. The median duration between onset of ventilation and colorimetric CO₂ detection was 14 (IQR 11–19) s. In three cases, the infant was not successfully intubated by 3 min after birth, and the umbilical cord was therefore clamped and cut prior to intubation. Two of these infants were born via vaginal delivery and one via caesarean delivery. Neonatal providers in those cases identified that infant positioning was the major impediment to procedural success, and all three of these infants were intubated after UCC once placed on the warmer bed.

Safety outcomes were similar between trial participants and historical controls (table 2). The first measured haemoglobin was higher among trial participants than historical controls (table 3). Mean blood pressure values were higher in trial participants at 1 hour of life. Subsequent mean blood pressure values assessed at 6 hours of life and beyond were similar between groups (data not shown). Blood gas parameters and oxygenation indices did not differ between trial participants or controls on the first blood gas or at any point in the first 48 hours after birth. There were no differences between groups in the use of vasoactive medications, inhaled nitric oxide or ECMO. No trial participants and one historical control died in the first week after birth.

Echocardiograms were obtained at a mean of 13 hours after birth. One trial participant and one historical control were supported with ECMO when the echocardiograms were obtained, and these images were not assessed. All infants had evidence of pulmonary hypertension (online supplementary table).
enrolled newborns. However, the protocol completion rate was lower among infants in the intervention group in that study (59%) compared with other trials of preterm infants (89%–100%).

Intubation is a standard approach to the delivery room management for all infants with CDH in our hospital. Thus, establishing lung aeration prior to UCC among infants with CDH requires intubation with an intact cord. In this trial, 17/20 infants were intubated within the specified 3 min time-frame before UCC. The threshold of allowing 3 min for intubation prior to UCC was considered to represent a reasonable balance between providing sufficient time for the neonatologist to intubate the infant without introducing an excessive delay in intubation and onset of ventilation if intubation before UCC was not possible. Neonatologists involved in unsuccessful cases expressed that positioning on the trolley was the major impediment to intubation, suggesting that additional time beyond 3 min may not have resulted in procedural success.

UCC was performed after colorimetric CO₂ detection. It is possible that a longer duration of ventilation after intubation would allow for more adequate lung aeration before UCC. Physiologically based cord clamping has been variably defined. Lefebvre et al specified for UCC to occur after ‘stabilisation’, indicated by measures of heart rate and oxygen saturation. Brouwer et al targeted vital sign parameters as well as exhaled tidal volume >4 mL/kg as a marker of lung aeration among preterm infants. This may not be a reasonable target for infants with CDH and related pulmonary hypoplasia, in whom tidal volumes after birth are often <4 mL/kg for both spontaneous breaths and manual inflations. In addition, tidal volume monitoring is not standard in most delivery rooms, and we sought to avoid introducing additional equipment into a crowded physical space for the study intervention.

Colorimetric CO₂ detectors are small, portable and standardly used to confirm intubation success. Expired CO₂ is correlated with tidal volume, and colorimetric CO₂ precedes rise in heart rate among bradycardic infants. Blank et al targeted UCC to occur ≥60 s after colorimetric CO₂ detection for infants who required respiratory support in a recent feasibility trial. In future studies of infants with CDH, it may be reasonable to provide ventilation for a similar duration after CO₂ detection before UCC.

This single-arm pilot trial was not designed to detect the impact of the study intervention on cardiopulmonary outcomes for infants with CDH. A randomised study powered for relevant outcomes is needed to address this question. There were transient differences in physiological outcomes such as haemoglobin and blood pressure between trial participants and historical controls. We assessed for echocardiographic evidence of pulmonary hypertension, but these studies—obtained for clinical indications—were performed on average 13 hours after birth and may not have reflected infants’ haemodynamic status immediately after birth. Lefebvre et al reported that infants treated with intact cord resuscitation experienced transiently higher systemic blood pressures and blood gas pH values. In the present study, blood gas parameters were similar between participants and historical controls at all assessed time points, and initial pH values were low in both groups. One potential explanation for this is that infants enrolled by Lefebvre et al had less severe antenatal parameters, with mean O/E LHR of 55%.

We acknowledge study limitations. Neonatal intubations were performed by attending neonatologists. Feasibility outcomes may not generalise to less experienced providers, who typically have lower intubation success rates. In addition, we did not

### Table 2 Maternal and infant safety outcomes

<table>
<thead>
<tr>
<th>Neonatal outcomes</th>
<th>Trial participants (n=20)</th>
<th>Historical controls* (n=19)</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cord avulsion</td>
<td>0</td>
<td>0</td>
<td>N/A</td>
</tr>
<tr>
<td>Chest compressions</td>
<td>0</td>
<td>0</td>
<td>N/A</td>
</tr>
<tr>
<td>Hypothermia on first temperature (&lt;36°C), n (%)</td>
<td>3 (15)</td>
<td>2 (11)</td>
<td>&gt;0.99</td>
</tr>
<tr>
<td>First temperature (°C), mean (SD)</td>
<td>36.7 (0.9)</td>
<td>36.8 (0.6)</td>
<td>0.56</td>
</tr>
</tbody>
</table>

### Table 3 Physiological outcomes

<table>
<thead>
<tr>
<th></th>
<th>Trial participants (n=19)</th>
<th>Historical controls (n=19)</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Apgar score at 1 min, median (IQR)</td>
<td>5 (3–7)</td>
<td>7 (3–8)</td>
<td>0.51</td>
</tr>
<tr>
<td>Apgar score at 5 min, median (IQR)</td>
<td>8 (5–8)</td>
<td>8 (5–9)</td>
<td>0.72</td>
</tr>
<tr>
<td>First haemoglobin, g/dL, mean (SD)</td>
<td>17.6 (1.3)</td>
<td>16.3 (1.9)</td>
<td>0.02</td>
</tr>
<tr>
<td>Mean blood pressure 1 hour after birth; mean (SD)*</td>
<td>51.1 (8.5)</td>
<td>44.3 (6.3)</td>
<td>0.008</td>
</tr>
<tr>
<td>First blood gas after birth*</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>pH, mean (SD)</td>
<td>7.02 (0.15)</td>
<td>7.03 (0.13)</td>
<td>0.74</td>
</tr>
<tr>
<td>CO₂, mean (SD)</td>
<td>90 (26)</td>
<td>88 (25)</td>
<td>0.82</td>
</tr>
<tr>
<td>Base deficit, (mean SD)</td>
<td>8.9 (3.3)</td>
<td>9.8 (3.8)</td>
<td>0.51</td>
</tr>
<tr>
<td>Oxygenation index with first blood gas, median (IQR)</td>
<td>17.5 (12.8–25.5)</td>
<td>16.3 (12.2–22.8)</td>
<td>0.74</td>
</tr>
<tr>
<td>Vasopressors (first 48 hours), n (%)</td>
<td>13 (68)</td>
<td>16 (84)</td>
<td>0.45</td>
</tr>
<tr>
<td>INO (first 48 hours), n (%)</td>
<td>9 (47)</td>
<td>11 (58)</td>
<td>0.52</td>
</tr>
<tr>
<td>ECMO (first 7 days), n (%)</td>
<td>7 (37)</td>
<td>4 (21)</td>
<td>0.48</td>
</tr>
<tr>
<td>Mortality (first 7 days), n (%)</td>
<td>0</td>
<td>1 (5)</td>
<td>&gt;0.99</td>
</tr>
</tbody>
</table>

*One trial participant was not matched due to postnatal diagnosis of major anomaly.

CO₂, cesarean section.

DISCUSSION

We conducted this pilot trial to assess the safety and feasibility of initiating resuscitation prior to UCC for infants with CDH. It was possible to position all enrolled infants on a trolley adjacent to their mothers after birth. In most cases, it was possible to perform immediate intubation and ventilation with an intact umbilical cord. We did not find any evidence that this approach to resuscitation poses increased risk to infants or their mothers. Enrolled infants had higher haemoglobin levels and transiently higher systemic blood pressures compared with matched historical controls.

This study contributes to a growing body of literature demonstrating the feasibility and safety of initiating resuscitation prior to UCC. In contrast to our study population, few infants in previous trials required invasive respiratory support. In the trial by Duley et al, more invasive interventions were performed, including intubation and cardiac resuscitation of...
capture granular data on physiological parameters such as heart rate and oxygen saturation during the initial resuscitation, and non-invasive measures of pulmonary blood flow are not readily available. Study strengths include a prospective trial design with extensive training and oversight by the study team. We used a physiology-based, rather than time-based, approach to UCC. 27 Finally, the trial included infants with a wide spectrum of prognostic CDH parameters, including severely affected infants.

In conclusion, performing intubation and initiating ventilation prior to UCC is safe and feasible among infants with CDH. The impact of this approach on clinically relevant outcomes deserves investigation in a randomized trial.

Acknowledgements The authors gratefully acknowledge Sara DeMauro, Lakshmi Srinivasan and Robert Sutton for serving on the Data Safety Monitoring Committee; Karen Pesta, Joel Stejskal, Alme Coursey and Leane Sookian for their assistance training clinical staff; and the Division of Neonatology for providing financial support for equipment used in this trial. The authors would like to thank all of the mothers and infants who participated in this trial and our clinical colleagues who supported the study.

Contributors The authors made the following contributions to this work: study design: EEF, AA, HLH, NR, DAM, JM, JG, BS, AC and SJR. Data collection and analysis: EEF, AC, DW, NN, MVF and SJR. Drafting Manuscript: EEF. Critically revise manuscript: all authors.

Funding EEF is supported by a National Institute of Child Health and Human Development Career Development Award, K23HD084727. NN receives research support from Draeger, Aerogen, Smiths Medical, Actuated Medical and Vero Biotech.

Competing interests EEF is an unpaid member of the Scientific Advisory board for Concord Neonatal. NN receives research support from Draeger, Aerogen, Smiths Medical, Actuated Medical and Vero Biotech. No other competing interests are declared.

Patient consent for publication Parental/guardian consent obtained.

Ethics approval This study was approved by the Children’s Hospital of Philadelphia Institutional Review Board (17–0 14 125).

Provenance and peer review Not commissioned; externally peer reviewed.

Data availability statement All data relevant to the study are included in the article or uploaded as supplementary information.

REFERENCES