Effect of Minimally Invasive Surfactant Therapy vs Sham Treatment on Death or Bronchopulmonary Dysplasia in Preterm Infants With Respiratory Distress Syndrome
The OPTIMIST-A Randomized Clinical Trial

Peter A. Dargaville, MD; C. Omar F. Kamlin, DMedSci; Francesca Orsini, MSc; Xiaofang Wang, PhD; Antonio G. De Paoli, MD; H. Gozte Kanmaz Kutman, MD; Merih Cetinkaya, PhD; Lilijana Kornhauser-Cerar, PhD; Matthew Derrick, MBBS; Hilal Özkan, MD; Christian V. Hulzebos, PhD; Georg M. Schmolzer, PhD; Ajit Aiyappan, PhD; Brigitte Lemyre, MD; Sheree Kuo, MD; Victor S. Rajadurai, MD; Joyce O’Shea, MD; Manoj Binivale, MD; Rangasamy Ramanathan, MD; Alla Kashmiri, MD; David Bader, MD; Mark R. Thomas, MD; Mallinath Chakraborty, PhD; Mariam J. Buksh, MD; Risha Bhatia, PhD; Carol L. Sullivan, MD; Eric S. Shinwell, MD; Amanda Dyson, MMEd; David P. Barker, DM; Amir Kugelman, MD; Tim J. Donovan, MPH; Markus K. Tauscher, MD; Vadivelam Murthy, MD; Sanoj K. M. Ali, MD; Pete Yossuck, MD; Howard W. Clark, DPhil; Roger F. Soll, MD; John B. Carlin, PhD; Peter G. Davis, MD; for the OPTIMIST-A Trial Investigators

IMPORTANCE The benefits of surfactant administration via a thin catheter (minimally invasive surfactant therapy [MIST]) in preterm infants with respiratory distress syndrome are uncertain.

OBJECTIVE To examine the effect of selective application of MIST at a low fraction of inspired oxygen threshold on survival without bronchopulmonary dysplasia (BPD).

DESIGN, SETTING, AND PARTICIPANTS Randomized clinical trial including 485 preterm infants with a gestational age of 25 to 28 weeks who were supported with continuous positive airway pressure (CPAP) and required a fraction of inspired oxygen of 0.30 or greater within 6 hours of birth. The trial was conducted at 33 tertiary-level neonatal intensive care units around the world, with blinding of the clinicians and outcome assessors. Enrollment took place between December 16, 2011, and March 26, 2020; follow-up was completed on December 2, 2020.

INTERVENTIONS Infants were randomized to the MIST group (n = 241) and received exogenous surfactant (200 mg/kg of poractant alfa) via a thin catheter or to the control group (n = 244) and received a sham (control) treatment; CPAP was continued thereafter in both groups unless specified intubation criteria were met.

MAIN OUTCOMES AND MEASURES The primary outcome was the composite of death or physiological BPD assessed at 36 weeks’ postmenstrual age. The components of the primary outcome (death prior to 36 weeks’ postmenstrual age and BPD at 36 weeks’ postmenstrual age) also were considered separately.

RESULTS Among the 485 infants randomized (median gestational age, 27.3 weeks; 241 [49.7%] female), all completed follow-up. Death or BPD occurred in 105 infants (43.6%) in the MIST group and 121 (49.6%) in the control group (risk difference [RD], −6.3% [95% CI, −14.2% to 1.6%]; relative risk [RR], 0.87 [95% CI, 0.74 to 1.03]; P = .10). Incidence of death before 36 weeks’ postmenstrual age did not differ significantly between groups (24 [10.0%] in MIST vs 19 [7.8%] in control; RD, 2.1% [95% CI, −3.6% to 7.8%]; RR, 1.27 [95% CI, 0.63 to 2.57]; P = .51), but incidence of BPD in survivors to 36 weeks’ postmenstrual age was lower in the MIST group (81/217 [37.3%] vs 102/225 [45.3%] in the control group; RD, −7.8% [95% CI, −14.9% to −0.7%]; RR, 0.83 [95% CI, 0.70 to 0.98]; P = .03). Serious adverse events occurred in 10.3% of infants in the MIST group and 11.1% in the control group.

CONCLUSIONS AND RELEVANCE Among preterm infants with respiratory distress syndrome supported with CPAP, minimally invasive surfactant therapy compared with sham (control) treatment did not significantly reduce the incidence of the composite outcome of death or bronchopulmonary dysplasia at 36 weeks’ postmenstrual age. However, given the statistical uncertainty reflected in the 95% CI, a clinically important effect cannot be excluded.

TRIAL REGISTRATION anzctr.org.au Identifier: ACTRN12611000916943

Published online December 13, 2021.
For the newly born preterm infant, published guidelines recommend continuous positive airway pressure (CPAP) for initial respiratory support rather than intubation and ventilation,\(^1,2\) with the expectation of equivalent or better outcomes, including a reduction in the risk of bronchopulmonary dysplasia, the chronic disease of the preterm lung.\(^3\) For the composite outcome of death or bronchopulmonary dysplasia, a meta-analysis of data from clinical trials\(^4-7\) in preterm infants with a gestational age of less than 30 weeks showed an association favoring initial CPAP with a relative risk (RR) of 0.91 (95% CI, 0.84-0.99).\(^8\) In these trials, exogenous surfactant, which is a proven therapy for respiratory distress syndrome (RDS),\(^9\) was only administered in the CPAP group if relatively high thresholds (0.40-0.60) for fraction of inspired oxygen (FIO\(_2\)) exceeded.

It remains unclear whether the benefit of initial CPAP on respiratory outcomes in these trials would have been greater if, in the CPAP group, surfactant had been selectively administered at a lower FIO\(_2\) threshold in infants with more prominent features of RDS. CPAP alone often fails to provide sufficient support in such cases,\(^10,11\) and in a large nonrandomized study\(^12\) this pathway of delayed intubation was associated with deleterious consequences, including an increased risk of death or bronchopulmonary dysplasia.

Without an endotracheal tube, preterm infants supported with CPAP lack the usual conduit for instillation of exogenous surfactant, thus invoking the dilemma of how to administer surfactant therapy to those with features of RDS.\(^9\) Among numerous less invasive alternatives, intratracheal administration of surfactant via a thin catheter has emerged as a practicable and propitious solution,\(^13-15\) but the benefits of this approach, including in relation to death or bronchopulmonary dysplasia, remain uncertain.

The OPTIMIST-A trial (one of cOllaborative Paired Trials Investigating Minimally Invasive Surfactant Therapy) tested the hypothesis that administration of surfactant via a thin catheter (minimally invasive surfactant therapy [MIST]) would reduce the incidence of the composite outcome of death or bronchopulmonary dysplasia or its components.

**Methods**

**Study Design and Oversight**

The trial was an investigator-initiated, international, multicenter randomized clinical trial of a blinded intervention conducted at 33 tertiary-level neonatal intensive care units in Australia, Canada, Israel, New Zealand, Qatar, Singapore, Slovenia, the Netherlands, Turkey, the UK, and the US. The human research ethics committees of all participating centers approved the trial protocol\(^16\) (Supplement 1). Prospective written parental consent was obtained prenatally or postnatally. An independent data and safety monitoring committee reviewed the interim analyses for safety and efficacy. The statistical analysis followed the statistical analysis plan (Supplement 2), in which the method of analysis outlined in the trial protocol was expanded upon, including nomination of the components of the primary outcome as end points, and identification of a group of key clinical and safety outcomes from among the secondary outcomes.

**Randomization and Masking**

Randomization and the intervention were undertaken by a treatment team comprising staff not currently involved in the infant’s care, including at least 1 proceduralist trained in the technique of MIST and an assistant (neonatal nurse or respiratory therapist).

Infants were randomized 1:1 to the MIST group or the sham treatment (control) group via a computer-generated code linked to a corresponding opaque sealed envelope (Figure). The randomization sequence used permuted block sizes of 2, 4, or 6 with stratification by study center and gestational age. Multiple births were randomized independently.

Clinicians and parents were blinded to the study intervention with screening of the infant’s bedspace from external view. Central physiological monitors were disconnected.
whenever possible and a study oximeter was used to display heart rate and oxygen saturation.

Interventions

The MIST intervention was administered using the Hobart method as described previously6,18 (Video; a longer version is also available at https://youtu.be/wAkNATfH9S0). Administration of a sedating premedication was not done; intranasal 25% sucrose, intravenous atropine, or both, were allowed. Surfactant was instilled using a 16-gauge vascular catheter (Angiocath, Becton Dickinson) or a proprietary catheter (LIStAcath, Chiesi Farmaceutici), marked to indicate the correct insertion depth beyond the vocal cords (1.5 cm for gestational age <27 weeks; 2 cm otherwise). Via direct laryngoscopy, the catheter was inserted into the trachea (maximum of 3 attempts) and surfactant (200 mg/kg of poractant alfa; Chiesi Farmaceutici) was administered in 3 to 4 aliquots, with a 10-second pause between each. CPAP was applied throughout, aiming to optimize surfactant dispersion from the trachea by spontaneous breathing. Infants with apnea initially received tactile stimulation. Positive pressure inflations by mask were used for refractory apnea, persistent hypoxemia, or bradycardia.

Infants in the control group received a sham intervention consisting only of transient repositioning. The interactions of the treatment team and the duration of the procedure mimicked that of the MIST procedure.

After the study intervention, infants were returned to their original position, standard monitoring was restored, and care assured by treating clinicians. Three members of the staff, including the bedside nurse, completed a brief questionnaire that asked whether they could discern which intervention the infant had received.

Infants in both groups were intubated if requiring FIO2 of 0.45 or greater (or by clinician discretion when requiring FIO2 >0.40) or if there was severe or recurrent apnea or persistent respiratory acidosis. Once intubated, surfactant could be administered according to clinical judgement. Surfactant administration by thin catheter was not allowed in either group due to the intervention. Other aspects of clinical management were not protocolized.

Outcomes

The primary outcome was the composite of death prior to 36 weeks’ postmenstrual age or physiological bronchopulmonary dysplasia assessed at 36 weeks’ postmenstrual age by blinded study personnel. Infants were assigned a diagnosis of bronchopulmonary dysplasia if at 36 weeks’ postmenstrual age they were: (I) supported with mechanical ventilation, CPAP, or high-flow nasal cannula therapy (at a rate of
≥2 L/min), or (2) receiving supplemental oxygen with actual or effective FIo2 of 0.30 or greater, or (3) receiving oxygen with FIo2 less than 0.30 and did not pass an air trial or no air trial was done per physician request.

The secondary outcomes included the components of the primary composite outcome and 6 key clinical and safety outcomes that were prespecified based on their importance in relation to early respiratory management, in-hospital care, and potential effect on long-term outcome. The outcomes were pneumothorax requiring drainage, need for intubation within 72 hours of birth, grade III or IV intraventricular hemorrhage, the composite of death during hospitalization or major morbidity, and each of these considered separately. Major morbidity was defined as any of the following: intraventricular hemorrhage grade III or IV; cystic periventricular leukomalacia; retinopathy of prematurity stage 3 or greater; or physiological bronchopulmonary dysplasia. There were 15 additional binary secondary outcomes and 7 continuous secondary outcomes.

Additional protocized secondary outcomes not reported herein include length of stay in the intensive care unit (variably defined worldwide), hospital billings and calculated cost of hospitalization, and outcomes beyond the first hospitalization, including death or major disability at 2 years’ postmenstrual age and respiratory morbidity during the first 2 years. Exploratory procedural and safety outcomes related to the MIST intervention were ascertained as were adverse events. Further details appear in the statistical analysis plan (Supplement 2). All deaths and unexpected serious adverse events were reported to local ethics committees and the trial management center within 5 days and were reviewed by the data and safety monitoring committee.

**Sample Size Calculation**

The study aimed to recruit 606 infants, providing 90% power to detect an absolute risk reduction of 13% (RR reduction of 33%) in the incidence of death or bronchopulmonary dysplasia, which for the control group was projected to be 38% based on observational data.

**Statistical Analysis**

For the primary outcome and its components, the RR (with 95% CI) comparing active treatment with control was estimated according to randomization group using a generalized

---

### Table 1. Baseline Characteristics

<table>
<thead>
<tr>
<th></th>
<th>Minimally invasive surfactant therapy (n = 241)*</th>
<th>Control treatment (n = 244)*</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Demographic</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Age, median (IQR), wk</td>
<td>2.9 (1.4)</td>
<td>2.8 (1.5)</td>
</tr>
<tr>
<td>Continuous positive airway pressure level required at randomization, median (IQR), cm H2O</td>
<td>7 (6-8)</td>
<td>7 (6-8)</td>
</tr>
<tr>
<td>Fraction of inspired oxygen</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Level required at randomization, median (IQR)</td>
<td>0.35 (0.30-0.39)</td>
<td>0.35 (0.30-0.40)</td>
</tr>
<tr>
<td>Level required ≤0.35</td>
<td>152 (63.1)</td>
<td>150 (61.5)</td>
</tr>
</tbody>
</table>

* Data are expressed as No. (%) unless otherwise indicated.

b Indicates success of transition at birth. The score range is 0 to 10. A score of 0 to 2 is given for each of the following: heart rate, respiratory effect, reflex irritability, muscle tone, and skin color. An Apgar score greater than 7 at 5 minutes after birth generally indicates a satisfactory transition for a preterm infant.
Table 2. Primary Outcome Analysis

<table>
<thead>
<tr>
<th>No./total (%)</th>
<th>Minimally invasive surfactant therapy</th>
<th>Control treatment</th>
<th>Risk difference, % (95% CI)</th>
<th>Relative risk (95% CI)</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Death or bronchopulmonary dysplasia(^a)</td>
<td>105/241 (43.6)</td>
<td>121/244 (49.6)</td>
<td>−6.1 (−14.2 to 1.6)</td>
<td>0.87 (0.74 to 1.03)</td>
<td>.10</td>
</tr>
<tr>
<td>Death prior to 36 weeks' postmenstrual age</td>
<td>24/241 (10.0)</td>
<td>19/244 (7.8)</td>
<td>2.1 (−3.6 to 7.8)</td>
<td>1.27 (0.63 to 2.57)</td>
<td>.51</td>
</tr>
<tr>
<td>Bronchopulmonary dysplasia in survivors to 36 weeks' postmenstrual age(^b)</td>
<td>81/217 (37.3)</td>
<td>102/225 (45.3)</td>
<td>−7.8 (−14.9 to −0.7)</td>
<td>0.83 (0.70 to 0.98)</td>
<td>.03</td>
</tr>
</tbody>
</table>

\(^a\) Adjusted for gestational strata.
\(^b\) Bronchopulmonary dysplasia (physiological definition) was assessed at 36 weeks' postmenstrual age and was defined as (1) requiring mechanical respiratory support (mechanical ventilation, continuous positive airway pressure, or high-flow nasal cannula therapy, \(≥2\) L/min); (2) if not requiring mechanical respiratory support, requiring oxygen with actual or effective fraction of inspired oxygen \((\text{FiO}_2)\) of 0.30 or greater; (3) receiving oxygen with \(\text{FiO}_2\) less than 0.30 and failed air trial; or (4) receiving oxygen with \(\text{FiO}_2\) less than 0.30 and air trial not performed per physician request. (n = 2 infants; 1 infant in each treatment group).

Results

Infants were enrolled between December 16, 2011, and March 26, 2020. Four planned data and safety monitoring committee interim analyses of safety and efficacy (blinded to the study group) recommended that recruitment continue. Recruitment ceased on March 26, 2020, after research activities were put in abeyance indefinitely at most participating centers due to the COVID-19 pandemic. The trial steering committee, unaware of the interim study results, ceased enrollment at this juncture with 488 infants recruited. Patient follow-up (to hospital discharge) was completed on December 2, 2020, and 1 infant in the MIST group remained in the hospital.

Study Infants

A total of 5187 infants were screened at 33 participating centers, 488 were randomized, and data from 485 infants were included in the primary analysis (Figure). These infants had a median gestational age of 27.3 weeks (IQR, 26.4-28.1 weeks) and 241 of 485 infants (49.7%) were female. Baseline characteristics of the study infants overall were similar between the groups (Table I); however, the frequency of male sex, incomplete or no steroid exposure, and multiple birth within the gestational age stratum of 25 to 26 weeks were each 12% to 14% higher in the MIST group (eTable 1 in Supplement 3).

Primary Outcome and Components of the Primary Outcome

Death or bronchopulmonary dysplasia assessed at 36 weeks' postmenstrual age occurred in 105 infants (43.6%) in the MIST group and in 121 infants (49.6%) in the control group (\(P = .10\)) (Table 2). Death occurred prior to 36 weeks' postmenstrual age in 24 infants (10.0%) in the MIST group and in 19 infants (7.8%) in the control group (\(P = .51\)) (Table 2). Death occurred prior to 36 weeks' postmenstrual age in 24 infants (10.0%) in the MIST group and in 19 infants (7.8%) in the control group (\(P = .51\)). The incidence of bronchopulmonary dysplasia in survivors to 36 weeks' postmenstrual age was reduced in the MIST group (81/217 [37.3%] vs 102/225 [45.3%]) in the control group; \(P < .001\). This pattern of findings for the primary outcome and components of the primary outcome was unchanged in the analysis using the extended GLM with additional covariates (eTable 2 in Supplement 3) and in the as-treated and per-protocol populations (eTable 3 in Supplement 3).

Secondary Outcomes

Among the 6 key clinical and safety outcomes, the need for intubation within 72 hours of birth was reduced in the MIST group (36.5% vs 72.1% in the control group; \(P = .001\)), as was the incidence of pneumothorax requiring drainage (4.6% vs 10.2%; \(P = .14\%)), RR, 0.44 [95% CI, 0.25 to 0.78]; (0.30-0.35 and >0.35), and geographic region, assessing any interaction between subgroup and treatment assignment in the GLM. The frequency of missing data was low and available case analysis was used. Two-tailed \(P\) values <.05 were labeled as significant.
The cumulative proportion of infants requiring intubation in the 2 groups diverged during the first 24 hours (eFigure 1 in Supplement 3). The incidence of other key clinical and safety outcomes based on their primary in relation to early respiratory management, in-hospital care, and potential effect on long-term outcome in the gestational age range in this study.

Significant differences favoring the MIST group were noted for 4 of 15 binary secondary outcomes (Table 4). Treatment with MIST was associated with a reduced requirement for intubation at any time (54.8% vs 81.1% in the control group; RD, −26.7% [95% CI, −39.8% to −13.5%]; RR, 0.67 [95% CI, 0.54 to 0.84]), incidence of patent ductus arteriosus requiring medical therapy (35.3% vs 45.5%; RD, −10.5% [95% CI, −20.2% to −0.9%]; RR, 0.77 [95% CI, 0.60 to 0.99]), and in need for oxygen therapy at home in survivors to hospital discharge (14.7% vs 21.9%; RD, −7.1% [95% CI, −11.6% to −2.5%]; RR, 0.68 [95% CI, 0.52 to 0.88]).

The requirement for surfactant therapy via endotracheal tube was reduced after treatment with MIST (32.8% vs 68.4% in the control group); the mean total number of doses of surfactant administered was 1.42 in the MIST group and 0.96 in the control group (Table 4). The MIST group showed reductions in 4 of 6 continuous secondary outcomes quantifying duration of therapy: mechanical ventilation (1 day vs 4 days in the control group; median difference, −1.96 [95% CI, −3.19 to −0.73] days), CPAP (17 days vs 22 days; median difference, −4.62 [95% CI, −8.41 to −0.84] days), mechanical ventilation plus CPAP (25 days vs 32 days; median difference, −8.13 [95% CI, −13.98 to −2.27] days), and all forms of mechanical respiratory support (40 days vs 45 days; median difference, −6.42 [95% CI, −11.95 to −0.89 days]) (Table 4).

**Subgroups**

In an exploratory analysis by gestational age subgroups for the outcome of death prior to 36 weeks’ postmenstrual age, there was a statistically significant interaction in relation to the treatment effect (control group favored at lower gestational age and MIST group favored at higher gestational age; *P* = .01 for interaction), but not for the composite of death or bronchopulmonary dysplasia or for bronchopulmonary dysplasia in survivors (eTable 4 in Supplement 3). Evidence of interaction between gestational age stratum and group allocation also was seen in relation to the need for intubation within 72 hours of birth and death during hospitalization (eTable 5 in Supplement 3). The need for intubation within 72 hours of birth was 49% for the MIST group vs 72% for the control group within the gestational age stratum of 25 to 26 weeks and was 29% vs 72% within the gestational age stratum of 27 to 28 weeks (*P* = .02 for interaction).

For the secondary outcomes, the pattern of treatment effects did not appear to differ between gestational age strata (eTable 6 in Supplement 3). Subgroup analysis by FiO₂ level required at randomization showed no statistically significant interaction between the subgroups (eTables 7-8 in Supplement 3). Incidence of the primary outcome and its components varied considerably in the control group between geographic regions (eFigure 2 in Supplement 3), but there was no statistically significant interaction in the effect of MIST across regions (*P* > .05 for interaction terms for the primary outcome and its components).

**Exploratory Outcomes**

The trachea was cannulated during the first attempt in 76% of the MIST interventions (eTable 9 in Supplement 3). Hypoxemia and bradycardia were common but transient, with positive pressure inflations applied in 14% of cases, and emergent intubation needed in 1 infant. The surfactant instillation procedure was found to be effective (defined in footnote for eTable 9 in Supplement 3) in 88% of cases, with a median absolute reduction of 0.10 in FiO₂ level required at 4 hours after the procedure. Clinical staff correctly identified which intervention had been performed in 33% of cases, were incorrect in 11%, and were unsure in 56%.

**Adverse Events**

A total of 54 serious adverse events were reported (eTable 10 in Supplement 3), which were evenly distributed between treatment groups (10.3% of infants in the MIST group vs 11.1% in the control group) and were not considered to be related to participation in the study. Infants who died were receiving comparable levels of respiratory support before the intervention with those who survived for both study groups. Within the MIST group, the frequency of procedural complications and

---

**Table 3. Prespecified Key Clinical and Safety Outcomes**

<table>
<thead>
<tr>
<th>Outcome</th>
<th>No./total (%)</th>
<th>Minimally invasive surfactant therapy</th>
<th>Control treatment</th>
<th>Risk difference, % (95% CI)</th>
<th>Relative risk (95% CI)</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pneumothorax requiring drainage</td>
<td>11/241 (4.6)</td>
<td>25/244 (10.2)</td>
<td>−5.8 (−10.2 to −1.4)</td>
<td>0.44 (0.25 to 0.78)</td>
<td>.005</td>
<td></td>
</tr>
<tr>
<td>Need for intubation within 72 h of birth</td>
<td>88/241 (36.5)</td>
<td>176/244 (72.1)</td>
<td>−35.8 (−47.2 to −24.4)</td>
<td>0.50 (0.40 to 0.64)</td>
<td>&lt;.001</td>
<td></td>
</tr>
<tr>
<td>Intraventricular hemorrhage grade III or IV</td>
<td>18/241 (7.5)</td>
<td>24/244 (9.8)</td>
<td>−2.4 (−6.3 to 1.5)</td>
<td>0.75 (0.48 to 1.19)</td>
<td>.23</td>
<td></td>
</tr>
<tr>
<td>Death or major morbidity during first hospitalization</td>
<td>116/241 (48.1)</td>
<td>136/244 (55.7)</td>
<td>−7.9 (−18.6 to 2.7)</td>
<td>0.86 (0.70 to 1.05)</td>
<td>.14</td>
<td></td>
</tr>
<tr>
<td>Death during first hospitalization</td>
<td>28/241 (11.6)</td>
<td>20/244 (8.2)</td>
<td>3.3 (−2.2 to 8.9)</td>
<td>1.41 (0.73 to 2.69)</td>
<td>.30</td>
<td></td>
</tr>
<tr>
<td>Major morbidity during first hospitalization in survivors</td>
<td>88/213 (41.3)</td>
<td>116/224 (51.8)</td>
<td>−10.3 (−20.8 to 0.2)</td>
<td>0.80 (0.64 to 1.00)</td>
<td>.05</td>
<td></td>
</tr>
</tbody>
</table>

*The outcomes are listed in likely chronological order of appearance and were prespecified as key clinical and safety outcomes based on their primary in relation to early respiratory management, in-hospital care, and potential effect on long-term outcome in the gestational age range in this study.*

*Major morbidity includes intraventricular hemorrhage grade III or IV, cystic periventricular leukomalacia, retinopathy of prematurity greater than stage II (assessed throughout hospitalization), or physiological bronchopulmonary dysplasia at 36 weeks’ postmenstrual age.*

---

**Variables:**
- The cumulative proportion of infants requiring intubation in the 2 groups diverged during the first 24 hours (eFigure 1 in Supplement 3).
- The need for intubation within 72 hours of birth was 49% for the MIST group vs 72% for the control group within the gestational age stratum of 25 to 26 weeks and was 29% vs 72% within the gestational age stratum of 27 to 28 weeks (*P* = .02 for interaction).
- For the secondary outcomes, the pattern of treatment effects did not appear to differ between gestational age strata (eTable 6 in Supplement 3).
- Subgroup analysis by FiO₂ level required at randomization showed no statistically significant interaction between the subgroups (eTables 7-8 in Supplement 3).
- Incidence of the primary outcome and its components varied considerably in the control group between geographic regions (eFigure 2 in Supplement 3), but there was no statistically significant interaction in the effect of MIST across regions (*P* > .05 for interaction terms for the primary outcome and its components).

**Exploratory Outcomes:**
- The trachea was cannulated during the first attempt in 76% of the MIST interventions (eTable 9 in Supplement 3).
- Hypoxemia and bradycardia were common but transient, with positive pressure inflations applied in 14% of cases, and emergent intubation needed in 1 infant.
- The surfactant instillation procedure was found to be effective (defined in footnote for eTable 9 in Supplement 3) in 88% of cases, with a median absolute reduction of 0.10 in FiO₂ level required at 4 hours after the procedure.
- Clinical staff correctly identified which intervention had been performed in 33% of cases, were incorrect in 11%, and were unsure in 56%.

**Adverse Events:**
- A total of 54 serious adverse events were reported (eTable 10 in Supplement 3), which were evenly distributed between treatment groups (10.3% of infants in the MIST group vs 11.1% in the control group) and were not considered to be related to participation in the study.
- Infants who died were receiving comparable levels of respiratory support before the intervention with those who survived for both study groups.
- Within the MIST group, the frequency of procedural complications and
the disease trajectory during the 24 hours after the procedure appeared to be similar (eTable 11 in Supplement 3).

**Discussion**

In this multicenter, randomized clinical trial in preterm infants supported with CPAP and exhibiting features of RDS, administration of surfactant via a thin catheter at a low oxygenation threshold, compared with continuation of CPAP, did not significantly reduce the incidence of the composite outcome of death or bronchopulmonary dysplasia at 36 weeks’ postmenstrual age. A clinically important true treatment effect cannot be excluded, with the 95% CI including the possibility of an absolute reduction in death or bronchopulmonary dysplasia of 14%, which was higher than originally hypothesized.

<table>
<thead>
<tr>
<th>Table 4. Secondary Outcomes</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>No./total (%)</strong>&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>Intubation at any time</td>
</tr>
<tr>
<td>Requirement for surfactant therapy via endotracheal tube</td>
</tr>
<tr>
<td>Pulmonary hemorrhage</td>
</tr>
<tr>
<td>Oxygen therapy at day 28 (in survivors to day 28)</td>
</tr>
<tr>
<td>Bronchopulmonary dysplasia&lt;sup&gt;c&lt;/sup&gt; in survivors to 36 weeks’ postmenstrual age</td>
</tr>
<tr>
<td>Mechanical respiratory support at 36 weeks’ postmenstrual age (in survivors)</td>
</tr>
<tr>
<td>Oxygen therapy at home in survivors to hospital discharge&lt;sup&gt;d&lt;/sup&gt;</td>
</tr>
</tbody>
</table>

**Respiratory binary outcomes**

**Nonrespiratory binary outcomes**

- Patent ductus arteriosus requiring medical therapy: 85/241 (35.3) vs 111/244 (45.5), −10.5 (−20.2 to 0.9), 0.77 (0.60 to 0.99).
- Late-onset sepsis: 55/241 (22.8) vs 63/244 (25.8), −3.2 (−13.7 to 7.3), 0.88 (0.58 to 1.33).
- Intraventricular hemorrhage any grade: 85/241 (35.3) vs 91/244 (37.3), −2.4 (−12.2 to 7.3), 0.93 (0.72 to 1.22).

**Continuous outcomes**

- No. of surfactant doses, median (IQR): (n = 241) 1 (1 to 2) vs (n = 244) 1 (0 to 1), −1.96 (−3.19 to −0.73)<sup>f,g</sup>.
- Duration of mechanical ventilation via endotracheal tube, median (IQR), d: (n = 240) 17 (6 to 33) vs (n = 244) 22 (8 to 35), −4.62 (−8.41 to −0.84)<sup>f,g</sup>.
- Duration of CPAP, median (IQR), d<sup>h</sup>: (n = 240) 25 (8 to 45) vs (n = 244) 32 (12 to 48), −8.13 (−13.98 to −2.27)<sup>f,a</sup>.
- Duration of all forms of mechanical respiratory support, median (IQR), d: (n = 240) 49 (14 to 60) vs (n = 244) 54 (25 to 64), −6.42 (−11.95 to −0.89)<sup>a</sup>.

Abbreviation: CPAP, continuous positive airway pressure.

<sup>a</sup> Unless otherwise indicated. The protocolized secondary outcomes not reported are: length of stay in intensive care unit (variably defined worldwide); hospital billings and calculated hospitalization cost; and outcomes beyond first hospitalization (additional details appear in Supplement 1).

<sup>b</sup> Adjusted for gestational strata.

<sup>c</sup> Clinically defined as the requirement for oxygen, mechanical respiratory support, or both, at 36 weeks’ postmenstrual age. Mechanical respiratory support included mechanical ventilation, CPAP, or high-flow nasal cannula therapy (at rate of ≥ 2 L/min).

<sup>d</sup> Data are missing for 2 infants in the minimally invasive surfactant therapy group.

<sup>e</sup> The mean number of doses given per infant was 1.42 in the minimally invasive surfactant therapy group and 0.96 in the control group.

<sup>f</sup> Data are severely skewed.

<sup>g</sup> Between-group difference expressed as median difference (95% CI).

<sup>h</sup> Includes noninvasive positive pressure ventilation.

<sup>i</sup> Between-group difference expressed as mean difference (95% CI).
Analysis of the components of the primary outcome showed an absolute increase in death of 2% prior to 36 weeks’ postmenstrual age favoring the control group (with an upper bound of 7.8%), which was not statistically significant. If representative of a real difference in mortality, this would clearly temper enthusiasm for MIST potentially arising from other findings of this trial. For both randomization groups, deaths were reported to be due to a multiplicity of causes and were distributed throughout the first months of life. Site investigators (who were blinded to treatment group) did not attribute any deaths to enrollment in the trial. In exploratory subgroup analysis for the outcome of death prior to 36 weeks’ postmenstrual age, there was a significant interaction suggesting a higher mortality risk in the gestational age stratum of 25 to 26 weeks associated with allocation to the MIST group. This finding may have been due in part to a chance imbalance in the risk profile for this subgroup, but behoves caution in application of MIST in preterm infants at the most immature gestational ages, particularly in regions in which mortality for such infants remains high.

There was an absolute decrease of 8% in bronchopulmonary dysplasia in survivors to 36 weeks’ postmenstrual age favoring the MIST group. An effect of MIST on the incidence of bronchopulmonary dysplasia has plausibility by virtue of the observed reduction in exposure to positive pressure ventilation by endotracheal tube, a known intermediary in the development of injury to the preterm lung.23,24 However, the pathogenesis of bronchopulmonary dysplasia is multifactorial,3 with many contributory factors not alterable by a single dose of surfactant on day 1.

One previous trial compared surfactant delivery via a thin catheter with continuation of CPAP in 220 preterm infants, noting a reduction in need for intubation within 72 hours of birth (RR, 0.61 [95% CI, 0.42-0.88]), but no clear effect on bronchopulmonary dysplasia at 36 weeks’ postmenstrual age (RR, 0.59 [95% CI, 0.26-1.36]).25 With a larger sample size, blinded intervention, and protocolized intubation thresholds, the present study provides more robust and definitive evidence regarding this mode of surfactant delivery.

This trial is, to our knowledge, the largest study to date to investigate administration of surfactant to preterm infants once an oxygen requirement of 30% has been surpassed. This is a lower threshold for selective surfactant therapy than used in previous large clinical trials in preterm infants with a gestational age of less than 30 weeks requiring CPAP, in which surfactant administration only occurred (after intubation) if the Fio2 requirement exceeded 0.40 to 0.60.4-7 The choice of an Fio2 entry threshold of 0.30 in the present study was based on data from a cohort of infants with gestational ages of 25 to 28 weeks who were receiving CPAP from the outset. Among infants for whom the Fio2 requirement exceeded 0.30 in early life, most had moderate or severe RDS radiologically, and there was a high rate of CPAP failure (odds ratio, 5.6 [95% CI, 1.7-18.0]).11 In the present study, exploratory subgroup analysis in infants with baseline Fio2 requirement in the range of 0.30 to 0.35 showed the pattern of treatment effects after MIST to be not discernibly different from those of infants with an Fio2 requirement greater than 0.35 at randomization, suggesting that an oxygen requirement of 30% (rather than higher) is an appropriate intervention threshold.

Given the design of the trial, the question of whether surfactant delivered via endotracheal tube at the same treatment threshold would have conferred the same benefits as MIST cannot be answered. However, a recent network meta-analysis of clinical trials including infants receiving noninvasive respiratory support identified surfactant delivery via a thin catheter, rather than by brief intubation, to be the strategy most strongly associated with a reduction in death or bronchopulmonary dysplasia.26 Furthermore, a Cochrane review of head-to-head trials of surfactant delivery via a thin catheter or endotracheal tube has provided strong evidence in favor of the thin catheter delivery method (for death or bronchopulmonary dysplasia: RR, 0.59 [95% CI, 0.48-0.73]).27

The Hobart method of MIST using a semirigid catheter appeared to be practicable, needing only 1 attempt in three-quarters of cases, in line with other studies.25-28,29 The proportion of infants experiencing transient bradycardia or hypoxemia was higher than reported previously, but in the case of hypoxemia the proportion was similar to standard intubation.30 The need for positive pressure ventilation by mask and for emergent intubation was low in the MIST group.

Limitations
This study has several limitations. First, after slow recruitment, enrollment ceased at the onset of the COVID-19 pandemic at 81% of the planned recruitment target, meaning that the study was underpowered for detection of a treatment effect of the magnitude originally hypothesized. The combination of the overall rate of commencement of CPAP (54%) at all study centers and the proportion of infants supported with CPAP that qualified for the trial (44%) imposed a barrier to recruitment. Lack of availability of treatment team personnel and loss of equipoise at some centers also played a role.

Second, data on numerous clinically relevant outcomes were collected and multiple statistical comparisons were performed, raising the possibility of type I error. A cautious interpretation of the findings is warranted.

Third, although the intervention was successfully blinded to the treating clinicians, members of treatment teams were likely to have participated in the care of enrolled infants at some time during hospitalization, raising the possibility of performance bias.

Conclusions
Among preterm infants with respiratory distress syndrome supported with CPAP, minimally invasive surfactant therapy compared with sham (control) treatment did not significantly reduce the incidence of the composite outcome of death or bronchopulmonary dysplasia at 36 weeks’ postmenstrual age. However, given the statistical uncertainty reflected in the 95% CI, a clinically important effect cannot be excluded.
Effect of Minimally Invasive Surfactant Therapy on Death or Bronchopulmonary Dysplasia in Preterm Infants

ARTICLE INFORMATION
Accepted for Publication: November 18, 2021.
Published Online: December 13, 2021.

Author Affiliations:
Menzies Institute for Medical Research, University of Tasmania, Hobart, Australia (Dargaville); Department of Paediatrics, Royal Hobart Hospital, Hobart, Australia (Dargaville, De Paoli); Neonatal Services, Royal Women’s Hospital, Melbourne, Australia (Kamlin, Davis); Department of Obstetrics and Gynaecology, University of Melbourne, Melbourne, Australia (Kamlin, Davis); Clinical Epidemiology and Biostatistics Unit, Murdoch Children’s Research Institute, Melbourne, Australia (Orsini, Wang, Carlin); Department of Neonatology, Zekiel Tahar Burak Maternity Teaching Hospital, Ankara, Turkey (Karmaz Kutman); Division of Neonatology, Department of Pediatrics, Istanbul Kanuni Sultan Suleyman Training and Research Hospital, Istanbul, Turkey (Cetinkaya); Department of Perinatology, Division of Gynaecology and Obstetrics, University Medical Centre, Ljubljana, Slovenia (Kornhauser-Cerar); Division of Neonatology, NorthShore University Health System, Evanston, Illinois (Derrick); Department of Pediatrics, Division of Neonatology, Uluďag University Faculty of Medicine, Bursa, Turkey (Özkân); Division of Neonatology, Beatrix Children’s Hospital, University Medical Center Groningen, Groningen, the Netherlands (Hulzebos); Division of Neonatology, Department of Pediatrics, University of Alberta, Edmonton, Canada (Schmölzer); Neonatal Services, Mercy Hospital for Women, Heidelberg, Australia (Aiyappan); Department of Obstetrics, Gynecology, and Newborn Care, Ottawa Hospital, Ottawa, Ontario, Canada (Lemyre); Department of Pediatrics, Kapiolani Medical Center for Women and Children, Honolulu, Hawaii (Kuo); Department of Neonatology, KK Women’s and Children’s Hospital, Duke-NUS Medical School, Singapore (Rajadurai); Neonatal Unit, Royal Hospital for Children, Glasgow, Scotland (Orsini); Division of Neonatology, Department of Pediatrics, LAC+USC Medical Center and Good Samaritan Hospital, Keck School of Medicine of USC, Los Angeles, California (Biniwale, Ramanathan); Department of Pediatrics, Children’s Regional Hospital, Cooper University Health Care, Camden, New Jersey (Kushnir); Department of Neonatology, Brai Zion Medical Center, Rappaport Faculty of Medicine, Technion-Israel Institute of Technology, Haifa, Israel (Bader); Department of Neonatal Medicine, Chelsea and Westminster Hospital NHS Foundation Trust, London, England (Domas); Regional Neonatal Intensive Care Unit, University Hospital of Wales, Cardiff (Chakraborty); Newborn Service, Starship Child Health, Auckland Hospital, Auckland, New Zealand (Buksh); Monash Newborn, Monash Children’s Hospital, Clayton, Australia (Bhatia); Department of Neonatology, Singleton Hospital, Swansea, Wales (Sullivan); Department of Neonatology, Ziv Medical Center, Faculty of Medicine, Bar-Ilan University, Tsfat, Israel (Shinwell); Department of Neonatology, Centenary Hospital for Women and Children, Canberra Hospital, Woden, Australia (Dysan); Neonatal Intensive Care Unit, Dunedin Hospital, Dunedin, New Zealand (Barker); Department of Neonatology, Rambam Medical Center, Rappaport Faculty of Medicine, Technion-Israel Institute of Technology, Haifa, Israel (Kugelman); Division of Neonatology, Royal Brisbane and Women’s Hospital, Brisbane, Australia (Donovan); Division of Neonatology, Peyton Manning Children’s Hospital, Ascension St Vincent, Indianapolis, Indiana (Tauscher); Neonatal Intensive Care Centre, Royal London Hospital-Barts Health NHS Foundation Trust, London, England (Murry); Division of Neonatology, Sir David Medical, Doha, Qatar (Al); Department of Pediatrics, WVU Medicine Children’s Hospital, Morgantown, West Virginia (Yossuck); Neonatal Intensive Care Unit, Princess Anne Hospital, Southampton, England (Clark); Division of Neonatal-Perinatal Medicine, Larner College of Medicine, University of Vermont, Burlington (Soll); Department of Paediatrics, University of Melbourne, Melbourne, Australia (Carlin).

Author Contributions: Dr Dargaville and Ms Orsini had full access to all of the data in the study and take responsibility for the integrity of the data and the accuracy of the data analysis.

Concept and design: Dargaville, Kamlin, Rajadurai, Soll, Carlin, Davis.


Drafting of the manuscript: Dargaville, Orsini, Soll, Carlin.

Critical revision of the manuscript for important intellectual content: Dargaville, Kamlin, Orsini, Wang, De Paoli, Kamraz Kutman, Cetinkaya, Kornhauser-Cerar, Derrick, Özkân, Hulzebos, Schmölzer, Aiyappan, Lemyre, Kuo, Rajadurai, O’Shea, Biniwale, Ramanathan, Kushnir, Bader, Thomas, Chakraborty, Buksh, Bhatia, Sullivan, Shinwell, Dyson, Barker, Kugelman, Donovan, Tauscher, Murthy, Ali, Yossuck, Clark, Carlin, Davis.

Data Sharing Statement: See Supplement 5.

Additional Contributions: We thank the families of infants in this study for their willingness to participate and are grateful for the assistance of staff at all sites in the conduct of the trial. We thank the members of the data and safety monitoring committee for their voluntary contribution to the trial: Brian A. Darlow, MD (committee chair; University of Otago), Michael Dunn, MD (University of Toronto), and Amy Salter, PhD (University of Adelaide). We thank the staff at the data management center at Murdoch Children’s Research Institute, including Ross Dunn, BAppSc, and Luke Stevens, BSc (Hons) (salaried data managers), and the staff at the trial coordinating center at the Menzies Institute for Medical Research, University of Tasmania, including Karen Butterley, RN, Nicky Stephens, MPhil, and Lizzy Reid, MSc (salaried trial coordinators).

Additional Information: The trial coordinating center and data management and statistics center were led by Drs Dargaville and Carlin, respectively.

REFERENCES
Effect of Minimally Invasive Surfactant Therapy on Death or Bronchopulmonary Dysplasia in Preterm Infants

Original Investigation Research


jama.com

© 2021 American Medical Association. All rights reserved.