Stem cells for bronchopulmonary dysplasia in preterm infants: A randomized controlled phase II trial

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Funding information
Korean Health and Medical Technology R&D Program, Ministry for Health, Welfare, and Family Affairs, Republic of Korea, Grant/Award Number: HI12C1821

Abstract
We previously demonstrated the safety and feasibility of mesenchymal stem cell (MSC) transplantation for bronchopulmonary dysplasia (BPD) in preterm infants in a phase I clinical trial. We thus investigated the therapeutic efficacy of MSCs for BPD in premature infants. A phase II double-blind, randomized, placebo-controlled clinical trial was conducted on preterm infants at 23 to 28 gestational weeks (GW) receiving mechanical ventilator support with respiratory deterioration between postnatal days 5 and 14. Infants were stratified by 23 to 24 GW and 25 to 28 GW and randomly allocated (1:1) to receive stem cells (1 × 10^7 cells/kg, n = 33) or placebo (n = 33). Although the inflammatory cytokines in the tracheal aspirate fluid were significantly reduced with MSCs, the primary outcome of death or severe/moderate BPD in the control group (18/33, 55%) was not significantly improved with MSC transplantation (17/33, 52%). In the subgroup analysis, the secondary outcome of severe BPD was significantly improved from 53% (8/15) to 19% (3/16) with MSC transplantation in the 23 to 24 GW group but not in the 25 to 28 GW subgroup. In summary, although MSC transplantation might be safe and feasible, this small study was underpowered to detect its therapeutic efficacy in preterm infants at 23 to 28 GW. Accordingly, we are now conducting an additional larger and controlled phase II clinical trial focusing on infants at 23 to 24 GW (NCT03392467). ClinicalTrials.gov identifier: NCT01828957.

KEYWORDS
bronchopulmonary dysplasia, cell transplantation, mesenchymal stem cells, premature infants

1 INTRODUCTION
Bronchopulmonary dysplasia (BPD) is a chronic lung disease that develops in premature infants receiving prolonged ventilator and oxygen therapy.1 Despite recent advances in neonatal intensive care medicine, BPD remains a major cause of mortality and long-term respiratory and neurologic morbidities in premature infants. A few clinically effective treatment options are available.2,3 New therapeutic strategies are, thus, urgently necessary to improve the prognosis of this serious intractable disorder.

Recently, our group and other research groups have reported that xenotransplantation of mesenchymal stem cells (MSCs) in immunocompetent newborn animals attenuates hyperoxic lung injuries such as impaired alveolarization, oxidative stress, inflammatory responses, increased apoptosis, and fibrosis, all of which simulate BPD in human infants.4-9 These protective effects of MSCs were cell type-dependent, showing better

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protection with umbilical cord blood (UCB)-derived MSCs than with adult adipose tissue- or bone marrow-derived MSCs \(^5,10\), time-dependent, showing better protection with early rather than late transplantation \(^6\); dose-dependent, showing better therapeutic efficacy with larger doses \(^7\); and route-dependent, showing better therapeutic efficacy with local intratracheal than systemic intraperitoneal or intravenous transplantation \(^8,11\). Furthermore, intratracheal transplantation of human UCB-derived MSCs was proven to be safe and feasible in recently conducted phase I clinical trials \(^12,13\) and was not associated with adverse respiratory, growth, or neurodevelopmental effects, as noted during the follow-up of these infants for up to 2 years \(^13\). However, therapeutic efficacy of this approach has not been assessed in a clinical setting to date. Here, we report a double-blind randomized placebo-controlled phase II clinical trial assessing the therapeutic efficacy of human UCB-derived MSC transplantation for BPD in extremely preterm infants.

### 2 | MATERIALS AND METHODS

#### 2.1 | Study design and participants

This phase II clinical trial was a randomized, double-blind, placebo-controlled two-center trial to assess the therapeutic efficacy of intratracheal transplantation of human UCB-derived MSCs for BPD in preterm infants conducted between 5 March 2013 and 23 March 2015, at the neonatal intensive care units (NICUs) of Samsung Medical Center (SMC) and Asan Medical Center (AMC), Seoul, Korea. The study protocol was approved by the Korean Food and Drug Administration (no. 11276) and by the institutional review boards (IRBs) of the SMC (IRB no. 2010-09-092) and AMC (IRB no. 2013-0333). External monitoring was independently performed by a contract research organization (Dream CIS Inc, Seoul, Korea). Written informed consent was obtained from both parents of each patient. A detailed study protocol was published at ClinicalTrials.gov (NCT01828957).

Preterm infants with a gestational age of 23 to 28 weeks, weighing 500 to 1250 g at birth, and who were on continuous invasive ventilator support because of respiratory deterioration were included in this study. Respiratory deterioration was defined as clinical signs of respiratory distress, including tachypnea and frequent desaturation (SaO\(_2\) < 80%), requiring an increase in ventilator settings within 24 hours before study enrollment during postnatal days 5 to 14. Patients were excluded for severe congenital anomalies, lung hypoplasia, severe septic shock, or severe (grade \(\geq 3\)) intraventricular hemorrhage (IVH). \(^14\)

#### 2.2 | Randomization

A total of 66 infants were enrolled in the trial; blocked randomization stratified according to gestational weeks (GW) (23-24 GW, \(n = 31\); 25-28 GW, \(n = 35\); Figure 1) was used. The MSC transplantation and control groups were assigned in the same ratio (1:1) in one block, and the randomization code table was prepared using SAS (SAS Institute Inc, Cary, North Carolina). Finally, 33 infants were assigned to the control group and 33 infants were assigned to the MSC group for analysis. The parents, hospital and research staff, and medical personnel directly involved in patient care were all blinded to group allocation throughout the study. The study drug was administered by an unblinded medical professional not involved in clinical management of the patients, and the administration was monitored by the other medical staff unaffiliated with the neonatal intensive care unit.

#### 2.3 | Procedures

For the treatment arm, standardized human UCB-derived MSCs at passage 6, prepared in strict compliance with good manufacturing practices (Pneumostem, Medipost Co, Seoul, Korea), were administered intratracheally via a gavage tube in two fractions into the left and right lungs at a total dose of \(1 \times 10^7\) cells/kg in 2 cc/kg of normal saline, as previously reported. \(^12\) An equal volume of normal saline (Medipost Co) was administered as the placebo to the control group.

#### 2.4 | Outcome measures

All patients were regularly assessed until 6 months after transplantation, and the clinical data were prospectively collected according to

### Lessons learned

- Intratracheal transplantation of human umbilical cord blood-derived mesenchymal stem cells in extremely preterm infants is safe and feasible.
- Because of the potential of stem cells for improving the incidence of severe bronchopulmonary dysplasia or death, especially in infants born at 23 to 24 gestational weeks, a study with larger sample size is required.

### Significance statement

A double-blind, randomized, placebo-controlled, phase II clinical trial of intratracheal transplantation of mesenchymal stem cells for bronchopulmonary dysplasia was performed for the first time. Stem cells showed a tendency, without statistical significance, to decrease the incidence of severe bronchopulmonary dysplasia or death, especially in infants at 23 to 24 gestational weeks, but not in those at 25 to 28 gestational weeks. Further study with a larger sample size would be required to solidly prove therapeutic benefits for preventing severe bronchopulmonary dysplasia or death in infants at 23 to 24 gestational weeks with the highest risk for bronchopulmonary dysplasia or death.
the schedule shown in the Supplemental Online Data. Clinical characteristics, including gestational age, birth weight, Apgar scores, sex, small size for gestational age, mode of delivery, antenatal steroid use, patent ductus arteriosus (PDA; defined as clinical signs of symptomatic PDA including heart murmur and wide pulse pressure plus ductal size >1.5 mm by echocardiography), and chorioamnionitis were analyzed. The tracheal aspirate fluid (TAF) was collected before transplantation and 7 days after MSC transplantation or placebo administration for the assessment of changes in the cytokine levels and factors known to be associated with the development or prevention of BPD. The samples were collected only when suctioning was clinically required during routine care; the details are described in the Supplemental Online Data.

Primary outcome measures included death or moderate/severe BPD, which was defined as the need for supplemental oxygen/respiratory support to maintain oxygen saturation >90% at 36 GW. Secondary outcome measures included mild, moderate, and severe BPD or death, pulmonary hemorrhage, duration of assisted respiratory support, oxygen use, hospitalization, postnatal steroid given with respiratory deterioration requiring increased ventilator settings, retinopathy of prematurity (ROP) stage ≥3 or requiring treatment, pneumothorax requiring treatment, pulmonary hemorrhage, growth, necrotizing colitis (stage >2b), blood culture proven nosocomial sepsis, and inflammatory cytokines in the TAF.

2.5 | Sample size and statistical analysis

The study sample size was determined for assessing the efficacy of MSCs compared with placebo with respect to the primary endpoint in order to decrease the incidence of moderate/severe BPD or death with a two-sided α level of 0.05% and 80% power. The target enrollment sample size of 70 and the minimum sample size of 62 necessary for assessing validation were calculated based on our previous phase I clinical trial in which a 38% decrease in the incidence of moderate/severe BPD or death was noted in the MSC transplantation group (33%) compared with that in a historical control group (71%). This is compatible with our nationwide Korean Neonatal Network data of the 74% prevalence of death or severe/moderate BPD in infants at 23 to 28 GW receiving invasive ventilator care for more than 14 days in 2013, and also accounted for a potential 10% dropout rate. The analysis plan for the primary endpoint was prespecified and performed with the per-protocol study population for testing efficacy. The data are expressed as means ± SD. The continuous variables were statistically compared between the groups using Student’s t test or the Mann-Whitney U test. The χ² test analysis was used for comparing the other nominal variables, and repeated measures analysis of variance was used for the assessment of the temporal profile of BPD biomarkers in the TAF. A value of P < .05 was considered statistically significant. The software package SPSS version 18 (IBM SPSS, Armonk, New York) was used for the aforementioned analyses. The Bayesian method was also used. To estimate the posterior probability of MSC treatment effect, that is, the relative risk (RR), 95% credible interval, and therapeutic benefit with the probability of a posterior RR of <1, Bayesian analysis was performed using SAS software version 14.3. R version 3.6.1 was used for Bayesian calculation of absolute risk reduction using the “neutral” prior of Laptook et al. The prior probability may be skeptical, neutral, or enthusiastic, as follows: (a) skeptical prior, assuming a 30% increase in the risk of death or BPD (RR, 1.30); (b) neutral prior, assuming no MSC treatment effect (RR, 1.0); and (c) enthusiastic prior, assuming a 37% decrease in the risk of death or BPD, where RR is estimated using binomial logistic regression analysis for analyzing the difference between severe BPD or death and moderate BPD and under.

3 | RESULTS

3.1 | Study population

Sixty-six infants were enrolled from a pool of 311 preterm infants born at 23 to 28 GW, with a birth weight of 500 to 1250 g, who born and admitted to SMC/AMC NICU between 5 March 2013 and 23 March 2015. Using a blocked randomization method, the enrolled
infants were stratified into subgroups based on gestational age (23-24 GW, n = 31; 25-28 GW, n = 35), and this study planned to recruit equal numbers of infants into each stratum. Enrolled infants were randomly allocated in a 1:1 ratio to either the MSC transplantation arm or the placebo control arm in each block (16 and 15 in the 23-24 GW subgroup and 17 and 18 in the 25-28 GW subgroup, respectively; Figure 1).

3.2 | Clinical characteristics

The demographical and clinical characteristics of the infants in each gestational age-stratified study group are shown in Table 1. Although the age at MSC transplantation showed a delayed tendency (P = .05) in the MSC group (12.1 ± 2.0 postnatal days) compared with the control group (10.3 ± 2.7 postnatal days) among the infants born at 23 to 24 GW, there were no significant differences in the other clinical characteristics, including gestational age, birth weight, and Apgar scores, between the study groups.

3.3 | Outcomes

The short-term outcomes of the infants in each gestational age-stratified study group are shown in Table 2. The primary outcome of death or severe/moderate BPD in the control group (18/33, 55%) was not significantly improved with MSC transplantation (17/33, 52%). In the subgroup analysis, the secondary outcome of severe BPD was significantly improved from 53% (8/15) to 19% (3/16) with MSC transplantation in the 23 to 24 GW group but not in the 25 to 28 GW subgroup. Other secondary outcomes, including duration of assisted respiratory support and hospitalization; incidence of ROP, pulmonary hemorrhage, and pneumothorax; steroid use for ventilator weaning; and incidence of other major complications associated with prematurity, were not significantly different between the study groups. No serious adverse events related to MSCs were observed until 6 months after transplantation. No newly developed abnormal mass lesions were found on chest radiography, and no significant abnormal results requiring correction were presented in laboratory tests, including complete blood count, glucose, aspartate aminotransferase (AST), alanine aminotransferase (ALT), total bilirubin, alkaline phosphatase, total protein, albumin, blood urea nitrogen, creatinine, and C-reactive protein, which were checked at 6 months after transplantation in all infants.

3.4 | Cause of death

Death before discharge was observed in 7% (1/15) of the patients in the control group and 19% (3/16) of the patients in the MSC group; this was noted among the infants born at 23 to 24 GW but not for those born at 25 to 28 GW (Table 2). One infant in the control group was extubated at postnatal day 56 and died of septic shock while on
TABLE 2  Short-term outcomes

| Outcome                        | Total  | MSC  | Control  | P value | MSC  | Control  | P value | Control  | P value |
|-------------------------------|--------|------|----------|---------|------|----------|---------|----------|---------|
| **Primary outcome**           |        |      |          |         |      |          |         |          |         |
| Moderate/severe BPD or death, n (%) | 18 (55) | 16 (49) | .62 | 11 (73) | 9 (56) | .32 | 7 (39) | 7 (41) | .89 |
| **Secondary outcomes**        |        |      |          |         |      |          |         |          |         |
| Mild BPD, n (%)               | 15 (46) | 17 (52) | .62 | 4 (27) | 7 (44) | .32 | 11 (61) | 10 (59) | .89 |
| Mod BPD, n (%)                | 3 (9) | 4 (12) | .69 | 2 (13) | 3 (19) | .68 | 1 (6) | 1 (6) | .97 |
| Severe BPD, n (%)             | 14 (42) | 9 (27) | .20 | 8 (53) | 3 (19) | .04 | 6 (33) | 6 (35) | .90 |
| Death at discharge, n (%)     | 1 (3) | 3 (9) | .30 | 1 (6) | 3 (19) | .32 | 0 (0) | 0 (0) | .89 |
| Severe BPD or death, n (%)    | 15 (46) | 12 (36) | .45 | 9 (60) | 6 (38) | .21 | 6 (33) | 6 (36) | .90 |
| **Respiratory support duration, median (IQR), days** |        |      |          |         |      |          |         |          |         |
| Intubation                    | 23 ± 17 | 24 ± 12 | .75 | 27 ± 12 | 24 ± 13 | .54 | 19 ± 19 | 24 ± 12 | .39 |
| Nasal CPAP                    | 36 ± 20 | 31 ± 15 | .26 | 39 ± 10 | 34 ± 11 | .27 | 34 ± 25 | 28 ± 17 | .48 |
| Total ventilator              | 59 ± 29 | 55 ± 19 | .55 | 66 ± 15 | 59 ± 21 | .27 | 53 ± 36 | 52 ± 18 | .97 |
| Oxygen duration               | 69 ± 35 | 68 ± 28 | .88 | 77 ± 22 | 71 ± 29 | .54 | 63 ± 42 | 66 ± 29 | .82 |
| Postnatal steroid, n (%)      | 23 (70) | 23 (70) | .73 | 11 (73) | 11 (69) | .78 | 12 (67) | 12 (71) | .80 |

R O P, n (%)

Grade ≥3                     | 13 (39) | 7/30 (23) | .17 | 8 (53) | 4/13 (31) | .23 | 5 (28) | 3 (18) | .48 |
| Treatment required           | 11/32 (34) | 8/30 (27) | .51 | 7/14 (50) | 4/13 (31) | .31 | 4 (22) | 4 (24) | .93 |
| Pneumothorax, n (%)          | 3 (9) | 1 (3) | .30 | 2 (13) | 1 (6) | .51 | 1 (6) | 0 (0) | .32 |
| Pneumonia, n (%)             | 8 (24) | 12 (36) | .28 | 4 (27) | 5 (31) | .78 | 4 (22) | 7 (41) | .23 |
| Pulmonary hypertension, n (%) | 1 (3) | 0 (0) | .31 | 0 (0) | 0 (0) | .89 | 1 (6) | 0 (0) | .32 |
| Pulmonary hemorrhage, n (%)  | 2 (6) | 1 (3) | .56 | 0 (0) | 0 (0) | .89 | 2 (11) | 1 (6) | .58 |
| Late-onset sepsis, n (%)     | 3 (9) | 6 (18) | .28 | 1 (7) | 3 (19) | .32 | 2 (11) | 3 (18) | .58 |
| NEC (stage ≥2b), n (%)       | 5 (15) | 3 (9) | .45 | 5 (33) | 1 (6) | .60 | 0 (0) | 2 (12) | .32 |
| Hospitalization duration, mean (SD), days | 107 ± 24 | 108 ± 28 | .91 | 108 ± 14 | 106 ± 30 | .80 | 107 ± 31 | 110 ± 26 | .75 |

Note: Data are presented as median ± interquartile range or as number and percentage of total.
Abbreviations: BPD, bronchopulmonary dysplasia; CPAP, continuous positive airway pressure; IQR, interquartile range; MSC, mesenchymal stem cell; NEC, necrotizing enterocolitis; ROP, retinopathy of prematurity.

TABLE 3  Bayesian analysis for mesenchymal stem cells benefit in reduction of death or BPD

| Outcome                        | Enthusiastic prior (RR, 0.74) | Neutral prior (RR, 1.0) | Skeptical prior (RR, 1.10) |
|-------------------------------|-----------------------------|-------------------------|---------------------------|
|                              | RR (95% CI) | P-TB, % | RR (95% CI) | P-TB, % | RR (95% CI) | P-TB, % |
| Moderate/severe BPD or death  |              |         |              |         |              |         |
| 23-24 weeks                   | 0.80 (0.42-1.21) | 84 | 0.83 (0.44-1.26) | 81 | 0.84 (0.46-1.29) | 78 |
| 25-28 weeks                   | 1.12 (0.49-1.91) | 42 | 1.20 (0.49-2.02) | 35 | 1.24 (0.52-2.08) | 31 |
| Total                         | 0.94 (0.55-1.35) | 64 | 0.97 (0.58-1.41) | 59 | 0.98 (0.58-1.42) | 56 |
| Severe BPD or death           |              |         |              |         |              |         |
| 23-24 weeks                   | 0.71 (0.29-1.16) | 89 | 0.75 (0.33-1.23) | 85 | 0.77 (0.34-1.28) | 83 |
| 25-28 weeks                   | 1.30 (0.35-2.51) | 55 | 1.09 (0.39-1.91) | 47 | 1.14 (0.42-2.03) | 42 |
| Total                         | 0.82 (0.42-1.28) | 79 | 0.86 (0.44-1.33) | 75 | 0.88 (0.45-1.36) | 72 |

Abbreviations: BPD, bronchopulmonary dysplasia; CI, Bayesian credible interval; P-TB, posterior probability of treatment benefit; RR, posterior risk ratio estimate.
continuous positive airway pressure (CPAP) support. Two infants in the MSC group also died of septic shock at postnatal days 26 and 59 while on mechanical ventilation. One infant in the MSC group was extubated at postnatal day 23 and died of intestinal perforation because of fulminant necrotizing colitis while on CPAP support.

### 3.5 Bayesian analyses

Bayesian analysis using a neutral prior indicated a 59% posterior probability of reduced incidence of death or moderate/severe BPD, with an estimated posterior risk ratio (PRR) of 0.97 (95% credible interval, 0.58-1.41) and 75% posterior probability of reduced incidence of death or severe BPD with an estimated PRR of 0.86 (95% credible interval, 0.44-1.33; Table 3). The posterior probability of reduced death or moderate/severe BPD was 64% and 56%, and probability of reduced death or severe BPD was 79% and 72%, respectively, when using enthusiastic prior or skeptical prior. Overall, the decrease in the incidence of severe BPD or death showed a higher probability of therapeutic benefit than a decrease in the incidence of moderate/severe BPD or death. In addition, this probability of therapeutic benefit was more prominent in the 23 to 24 GW group than in the 25 to 28 GW group.

Expressed as an absolute risk difference, using a neutral prior, the posterior probability that death or severe BPD in MSC group was at least 1%, 2%, or 3% less than control group was 70%, 67%, and 63%, respectively. A 2% absolute risk difference was associated with a 2.4-fold (67%/27%) higher probability of reduced, compared with increased death or severe BPD among MSC group infants relative to control group infants, assuming a range of risk differences viewed as equivalent.

### 3.6 Temporal profiles of cytokines and growth factors from TAF

The temporal profiles of cytokines and growth factors from the TAF are shown in Figure 2. Whereas the levels of the cytokines and growth factors at baseline showed no significant differences between the study groups, the levels of inflammatory cytokines, such as interleukin (IL)-1β, IL-6, IL-8, tumor necrosis factor-α, and matrix metalloproteinase-9, at post-transplantation day 7 were significantly lower in the MSC group than in the control group.

### 4 DISCUSSION

We performed a double-blind randomized placebo-controlled phase II clinical trial to assess the safety and therapeutic efficacy of MSC transplantation for BPD in extremely premature infants. We were unable to find previous randomized human studies evaluating the therapeutic efficacy of MSC transplantation for BPD in preterm infants.

In the present study, although intratracheal transplantation of MSCs in preterm infants was safe and feasible, this small study was underpowered to detect its statistically significant therapeutic efficacy for improving the primary outcome of death or severe/mild BPD in preterm infants at 23 to 28 GW. The underpower of this study might be primarily attributable to the much lower 32% rate of the primary outcome (especially in infants at 25-28 GW) than the 71% expected rate of the baseline primary outcome in the control group, and to the implausibly large 38% treatment effect size used for calculating the sample size of this trial. Nonetheless, considering the paucity of available clinical data on the therapeutic efficacy of MSC
transplantation, the reduced tendency of the primary outcome from 73% to 56% with MSC transplantation observed in the 23 to 24 GW subgroup may provide valuable information for future clinical trials with a larger sample size.

In this study, the gestational age cutoffs of 23 to 24 GW and 25 to 28 GW were used for the strata based on our previous institutional and nationwide Korean Neonatal Network data showing significant differences in mortality and the prevalence of at least moderate BPD especially between the 23 to 24 GW and 25 to 28 GW subgroups.\(^{19,20}\) The effect size of Pearson's correlation was \(-0.331\), indicating medium strength of the relationship between gestational age and the primary outcome of moderate/severe BPD or death. In subgroup analysis, the secondary outcome of severe BPD was significantly improved from 53% (8/15) to 19% (3/16) with MSC transplantation in the 23 to 24 GW group but not in the 25 to 28 GW subgroup. The Bayesian analysis using a neutral prior indicated an 81% posterior probability of reduced primary outcomes with stem cells relative to the placebo in the 23 to 24 GW subgroup. Accordingly, we are now conducting an additional larger controlled phase II clinical trial (NCT03392467) focusing on the most immature infants born at 23 to 24 GW who are at the highest risk of BPD or death.

Although male sex showed a tendency to be higher in the control group than in the MSC group, it was not associated with increased risk of BPD on multivariate analysis. The higher death rate in the MSC group than in the control group occurred only in the 23 to 24 and not in the 25 to 28 GW subgroup. This rate is much lower than the death group than in the control group, it was not associated with increased

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Death or BPD are difficult to predict in the first week of an infant’s life;\(^{26}\) hence, biomarkers might serve to aid early prediction of adverse outcomes beyond the rather crude tools of gestational age and the need for ventilation.\(^{20}\) Previously, using an animal model, we demonstrated that inflammation mediated by proinflammatory cytokines played a seminal role in the development of BPD\(^{4,6-8}\) and that the protection offered by MSCs against neonatal hyperoxic lung injury was primarily mediated by their paracrine anti-inflammatory, antioxidative, and antifibrotic effects rather than by the regenerative capacity of the MSCs.\(^{4,6-8}\) Furthermore, in concordance with our phase I clinical trial,\(^{12}\) the levels of inflammatory cytokines, such as IL-1β, IL-6, and IL-8, tumor necrosis factor-α, and matrix metalloproteinase-9, in the TAF were significantly reduced on post-MSC transplantation day 7 compared with those in the control group in this study. However, owing to our small sample size and lack of correlation with the clinical outcomes, our results pertaining to the biomarkers should be interpreted cautiously. Further studies are warranted to clarify whether any biomarker in the TAF could be used to predict death, severe BPD, and therapeutic efficacy of the MSCs.

Choosing the optimal MSCs for transplantation is a critical issue for successful clinical translation of stem cell therapy in preterm infants. UCB-derived MSCs exhibit several advantages over adult tissue-derived MSCs, including lower immunogenicity,\(^{10,27}\) higher proliferation capacity, paracrine potency, and therapeutic efficacy both in vitro\(^{5,28,29}\) and in vivo.\(^{5}\) Moreover, allogenic transplantation of MSCs might have a logistic advantage as they are ready to use “off the shelf” in the clinical setting. In the present study, we accordingly used the same clinical grade, standardized allogenic human UCB-derived MSCs, manufactured in strict compliance with the criteria for good manufacturing practice, from passage 6, as those used in previous phase I clinical trials for BPD\(^{12}\) and severe IVH.\(^{30}\)

Determining the optimal route for MSC transplantation is another important consideration. We previously demonstrated that the efficacy of local intratracheal transplantation of the MSCs is fourfold higher than that of systemic intravenous or intraperitoneal administration for protecting against hyperoxic lung injury in newborn rats, although the two routes are not mutually exclusive.\(^{8,11}\) In the present and previous studies,\(^{12}\) the MSCs were transplanted intratracheally in two fractions using the same method as that used for administering the exogenous surfactant, without any clinical instability or complications associated with MSC transplantation, suggesting that intratracheal administration of MSCs is safe and feasible. For intubated preterm infants receiving mechanical ventilation, additional procedures are not required for local administration. However, in a meta-analysis of MSCs in BPD, Augustine et al suggested that the intravenous route might be more efficacious than intratracheal administration of MSCs.\(^{31}\) Further studies are necessary to clarify this issue. Moreover, considering the recent increase in the use of noninvasive ventilation and less invasive techniques for surfactant administration,\(^{32}\) the feasibility of a new mode of MSC administration without intubation is another important issue that needs to be addressed in future clinical trials.

The major limitations of the present study include the small sample size, which was inadequate to attain statistically significant results. This was primarily due to the recently reduced incidence of BPD or death, even in the 25 to 28 GW subgroup of the control group, owing to improvements in neonatal intensive care medicine. These improvements limited the power of the study to detect small but important differences in the therapeutic efficacy of MSC transplantation according to the stratified GW subgroups, although the sample size was calculated appropriately. Accordingly, we are now conducting an additional larger and controlled phase II clinical trial (NCT03392467) focusing instead on the most immature infants born at 23 to 24 GW, who are at the highest risk of developing BPD or of death.
Nevertheless, the prospective double-blind randomized placebo-controlled design of the trial may be a strength of this phase II clinical trial. Another limitation might be the ability to detect safety concerns when fewer than 40 infants received the intervention. Other limitations include the lack of long-term respiratory and developmental outcome data. As we have just completed the 5-year long-term follow-up of these infants, we hope to publish these data in the near future.

5 | CONCLUSION

Although intratracheal transplantation of MSCs might be safe and feasible, it did not significantly improve the primary outcome of death or severe/moderate BPD in preterm infants at 23 to 28 GW. In subgroup analysis, the secondary outcome of severe BPD was significantly improved with MSC transplantation in the 23 to 24 GW group but not in the 25 to 28 GW subgroup. Accordingly, we are now conducting an additional longer and controlled phase II clinical trial, focusing on the most immature infants at 23 to 24 GW, who are at the highest risk of BPD or death (NCT03392467).

ACKNOWLEDGMENTS

We are grateful to Myoung Eun Lee, Clinical Research Coordinator, for her sincere assistance in administrative support, data collection, and management. This work was funded by a grant from the Korean Health and Medical Technology R&D Program, Ministry for Health, Welfare, and Family Affairs, Republic of Korea (HI12C1821). Human umbilical cord blood-derived mesenchymal stem cells were supplied by Medipost Co, Ltd. The sponsor had no involvement in study design, the collection, analysis, or interpretation of data, writing of the report, or the decision to submit the manuscript for publication.

CONFLICT OF INTEREST

Samsung Medical Center and Medipost Co, Ltd, have issued or filed patents for “Method of treating lung diseases using cells separated or proliferated from umbilical cord blood” under the names of Y.S.C. and W.S.P. The relevant application number is PCT/KR2007/000535. The other authors indicated no potential conflicts of interest.

AUTHOR CONTRIBUTIONS

S.Y.A., Y.S.C., A.-R.K., W.S.P.: conception/design, provision of study material or patients, data analysis and interpretation, manuscript writing, final approval of manuscript; M.H.L., S.I.S., B.S.L., K.S.K.: administrative support, collection and/or assembly of data, data analysis interpretation, manuscript writing, final approval of manuscript.

DATA AVAILABILITY STATEMENT

The data that support the findings of this study are available on request from the corresponding author. The data are not publicly available because of privacy or ethical restrictions.

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**SUPPORTING INFORMATION**

Additional supporting information may be found online in the Supporting Information section at the end of this article.

**How to cite this article:** Ahn SY, Chang YS, Lee MH, et al. Stem cells for bronchopulmonary dysplasia in preterm infants: A randomized controlled phase II trial. *STEM CELLS Transl Med*. 2021;10:1129–1137. [https://doi.org/10.1002/sctm.20-0330](https://doi.org/10.1002/sctm.20-0330)