Comparison of the Mortality and In-Hospital Outcomes of Preterm Infants Treated with Ibuprofen for Patent Ductus Arteriosus with or without Clinical Symptoms Attributable to the Patent Ductus Arteriosus at the Time of Ibuprofen Treatment

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Received: 7 June 2016
Accepted: 9 September 2016

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INTRODUCTION

Patent ductus arteriosus (PDA) is common in extremely preterm infants, affecting 60%–70% of infants with less than 28 weeks of gestation (1,2). While the ductus arteriosus remains open, blood flows left-to-right from the aorta into the pulmonary arteries, causing pulmonary edema, respiratory failure, and compromised perfusion of the bowel, kidney, and brain. There are strong epidemiologic evidences supporting that PDA is associated with numerous morbidities of preterm infants, such as intraventricular hemorrhage (IVH), necrotizing enterocolitis (NEC), retinopathy of prematurity (ROP), and bronchopulmonary dysplasia (BPD) (3-6). However, the extent to which these adverse outcomes are attributable to the hemodynamic consequences of ductal opening has not been established, and neither the best tools nor optimal thresholds to identify infants at greater risk for adverse outcomes have been delineated. Additionally, there is no consensus about the optimal timing of PDA treatment (7,8).

The aim of this study was to assess the differences in the mortality and in-hospital outcomes of preterm infants with < 28 weeks of gestation who received ibuprofen treatment according to the presence of clinical symptoms (any of oliguria, hypotension, or moderate to severe respiratory difficulty) attributable to hemodynamically-significant patent ductus arteriosus (hsPDA) at the time of first ibuprofen treatment. In total, 91 infants born from April 2010 to March 2015 were included. Fourteen infants (15.4%) received ibuprofen treatment when there were clinical symptoms due to hsPDA (clinical symptoms group). In clinical symptoms group, infants were younger (25 [23–27] vs. 26 [23–27] weeks; P = 0.012) and lighter (655 [500–930] vs. 880 [370–1,780] grams; P < 0.001). Also, the clinical risk index for babies (CRIB-II) scores were higher and more infants received invasive ventilator care ≤ 2 postnatal days. More infants received multiple courses of ibuprofen in clinical symptoms group. Although the frequency of secondary patent ductus arteriosus (PDA) ligation and the incidence of bronchopulmonary dysplasia (BPD) was higher in the clinical symptoms group in the univariate analysis, after multivariate logistic regression analysis adjusting for the CRIB-II score, birthweight, birth year, and the invasive ventilator care ≤ 2 postnatal days, there were no significant differences in mortality, frequency of secondary ligation and in-hospital outcomes including necrotizing enterocolitis (NEC), intraventricular hemorrhage (IVH), BPD or death. Our data suggest that we can hold off on PDA treatment until the clinical symptoms become prominent.

Keywords: Patent Ductus Arteriosus; Premature Infants; Treatment; Ibuprofen; Patient Outcome Assessment
Sehgal’s classification and the neonatal outcomes (13). However, the clinical criteria used in the McNamara staging system is non-specific and there are no studies searching for the efficacy of PDA treatment on neonatal outcomes according to the presence or severity of clinical symptoms.

Until the year 2013, our unit performed ibuprofen treatment for a PDA with a transductal diameter ≥ 1.4 mm/kg adapting the proposal by Kluckow and Evans (11). From 2013, our unit has adopted new strategies for the treatment guideline of hsPDA that are more focused on the presence of clinical symptoms as well as on the echocardiographic and radiographic results. We still do not know whether such guidelines are adequate in improving the outcomes of preterm infants especially associated with hsPDA.

The aim of this study was to assess whether there are any differences in the mortality and the in-hospital outcomes of preterm infants with a gestational age (GA) < 28 weeks who received ibuprofen treatment according to the presence of clinical symptoms attributable to the hsPDA at the time of first ibuprofen treatment. We also assessed the association between the echocardiographic and radiographic findings and the clinical symptoms due to hsPDA. And we assessed whether there was any difference in the pharmacologic characteristics of the ibuprofen treatment according to the presence of clinical symptoms attributable to the hsPDA.

MATERIALS AND METHODS

Study design

A retrospective study was done with 98 preterm infants < 28 weeks of gestation who were born and admitted to the neonatal intensive care unit of the Seoul National University Children’s Hospital between April 2010 and March 2015 and received ibuprofen for the treatment of hsPDA. Clinical and demographic data were collected from the reviewed medical records of the enrolled patients. Three infants were excluded because of a major congenital anomaly or the absence of echocardiographic data or information about the PDA treatment. We also excluded the infants with pulmonary hemorrhage, IVH ≥ grade 3, or NEC ≥ stage 2b before ibuprofen treatment because these morbidities are relative contra-indications for ibuprofen treatment. Finally, a total of 91 preterm infants were included in the analysis (Fig. 1).

If there is any suspect of cardiomegaly or pulmonary edema on chest radiograph, or respiratory difficulty, hypotension, decreased urine output or metabolic acidosis suggesting hsPDA, we checked the echocardiography for the confirmation of hsPDA. Before the ibuprofen treatment, all infants underwent an echocardiography to check for the presence of congenital heart defects and to evaluate the PDA status by a pediatric cardiologist. All infants received intravenous or oral ibuprofen at an initial dose of 10 mg/kg, followed by two doses of 5 mg/kg each after 24 and 48 hours. An oral route was preferred when feeding was started and there was no feeding intolerance. The patient was assessed clinically and echocardiographically at 24–48 hours after the third dose, and if the hsPDA persisted without any contraindications of ibuprofen treatment, a second and, if necessary, a third course of ibuprofen treatment was administered. If there were still significant respiratory difficulty, hypotension, or oliguria without closure of hsPDA by ibuprofen treatment, secondary PDA ligation was done.

We categorized the entire study population into two groups according to the presence of clinical symptoms attributable to the PDA at the time of first ibuprofen treatment. Then, we compared the baseline demographic characteristics prior to ibuprofen treatment including the GA at birth, birthweight, year of birth, gender, cesarean section, multiple pregnancy, complete course of antenatal steroid use, small for gestational age (SGA), clinical risk index for babies (CRIB)-II score (14), Apgar score at 5 minutes, presence of delivery room (DR) resuscitation, histo-

Fig. 1. Flow chart of the study population. A total of 91 infants with < 28 weeks of gestation born between April 2010 and March 2015 with hemodynamically-significant patent ductus arteriosus (hsPDA) receiving ibuprofen treatment were included in the analysis. Infants with hsPDA without clinical symptoms were 77 infants (84.6%) and infants with hsPDA with clinical symptoms were 14 infants (15.4%) at the time of first ibuprofen treatment.
logic chorioamnionitis, pregnancy induced hypertension (PIH), sepsis and inhaled nitric oxide use prior to treatment between the two groups. We examined the differences in echocardiographic findings just before the ibuprofen treatment and the presence of a causative organism documented from the blood sample use before ibuprofen treatment. Sepsis was defined as the hypotension with any inotropic or hydrocortisone use before ibuprofen treatment. Sepsis was defined as the presence of a causative organism documented from the blood culture. PDA closure rate was defined as the rate of PDA closure confirmed by echocardiography after ibuprofen treatment or secondary ligation. Secondary ligation was defined when infants received PDA ligation after failure to close the PDA with ibuprofen treatment. The postnatal day of PDA closure was defined as the first postnatal day of PDA closure confirmed by echocardiography in patients treated with only ibuprofen administration or the postnatal day of PDA ligation in infants receiving secondary ligation. We defined the duration of invasive ventilator care as the duration of ventilator support via endotracheal tube, and we also defined the duration of respiratory support as the whole duration of any ventilator or oxygen support during hospitalization.

Definitions
hsPDA was defined as a PDA with a transuductal diameter ≥ 1.4 mm/kg with significant left to right shunt confirmed by echocardiography considering the previous studies by Kluckow and Evans (11) and McNamara and Sehgal (12). Presence of clinical symptoms due to hsPDA was defined as at least one of the 3 following clinical symptoms besides echocardiographic confirmation compatible with the hsPDA:oliguria ≤ 1 mL/kg/hr in the preceding 8 hours before ibuprofen treatment, hypotension caused by PDA requiring inotropics more than 10 µg/kg/min or respiratory difficulty requiring invasive mechanical ventilator care. Respiratory difficulty was defined as having at least two of the following findings in infants with conventional ventilator care; FiO₂ > 0.5, respiratory rate > 40/min, and peak inspiratory pressure > 20 cmH₂O to maintain oxygen saturation between 88% and 95%, and a PaCO₂ < 65 mmHg. In infants with high frequency ventilator care, a mean airway pressure > 13 mmHg and a FiO₂ > 0.5 were required to define respiratory difficulty as the previous report (15).

First day of echocardiography was defined as the difference between the date of the first echocardiography examination and the date of birth. Zero day in the data meant the echocardiogram was done less than 24 hours after birth. First day of ibuprofen administration was determined from the difference between the date of first ibuprofen use and the date of birth. Cardiomegaly on the chest radiograph was defined as a cardiothoracic ratio of more than 0.6. The definition of SGA was according to the definition published by Olsen et al. (16). PIH was defined as any maternal diagnosis of eclampsia or preeclampsia. DR resuscitation was defined as cardiac compression or administration of medication in the DR. Invasive ventilator care ≤ 2 days after birth was defined as any ventilator support via endotracheal tube ≤ 2 days after birth. Hypotension before treatment was defined as hypotension with any inotropic or hydrocortisone use before ibuprofen treatment. Sepsis was defined as the whole duration of any ventilator or oxygen support during hospitalization.

Statistical analysis
All the continuous variables are expressed as the median (range), and the categorical variables are expressed as numbers and proportions. For univariate analysis, continuous variables were compared using the Mann-Whitney U test, and categorical variables were compared using the χ² or fisher’s exact test. Univariate analyses were done for the mortality, secondary PDA ligation, and in-hospital outcomes associated with prematurity including NEC ≥ stage 2b, spontaneous intestinal perforation (SIP), duration of respiratory support, duration of invasive ventilator care, hospital stay (days), moderate to severe BPD and severe BPD only. To assess the independent association between the presence of clinical symptoms due to hsPDA at the time of the first ibuprofen treatment and the primary outcomes (mortality ≥ 3 postnatal days, secondary PDA ligation and composite outcome including mortality ≥ 3 postnatal days, NEC ≥ stage 2b of modified Bell’s criteria [17], severe BPD at 36 weeks of gestation (18), and IVH ≥ grade 3 of Papile’s classification [19]) and secondary outcomes (NEC ≥ stage 2 or death, BPD or death, severe BPD or death, and IVH ≥ grade 3 or death), binary logistic regression analysis was done adjusting for the CRIB-II score, birthweight, the year of birth, and the invasive ventilator care as the duration of ventilator support via endotracheal tube, and we also defined the duration of respiratory support as ≤ 2 postnatal days. The statistical analysis was done with IBM SPSS Statistics version 20 (IBM Corp., Amarak, NY, USA) and R version 3.1.2. (http://www.r-project.org). P values less than 0.05 were considered statistically significant.

Ethics statement
This study was approved by the Institutional Review Board of Seoul National University Boramae Medical Center (IRB No. 16-2015-99). The need for informed consent was not required because of the retrospective nature of this study.

RESULTS
A total of 91 infants were included in our analysis. Seventy-seven infants (84.6%) received ibuprofen treatment for hsPDA without clinical symptoms at the time of first ibuprofen treatment.
and 14 infants (15.4%) were treated with clinical symptoms attributable to the hsPDA.

**Baseline and demographic characteristics**

Infants treated with ibuprofen in the presence of clinical symptoms due to hsPDA (clinical symptoms group) were younger and lighter at birth with higher CRIB-II scores compared to the infants without clinical symptoms due to hsPDA at the time of first ibuprofen treatment (no clinical symptoms group). Those infants who were born after 2013 tended to be more in the clinical symptoms group. More infants received invasive ventilator care within 2 postnatal days in the clinical symptoms group (Table 1).

**Echocardiographic and radiographic characteristics**

There was no statistically significant difference in the first day echocardiography between two groups [1 [0–7] day in no clinical symptoms group vs. 1 [0–3] day in the clinical symptoms group; \( P \) value = 0.803). Among the echocardiographic parameters at the time of first ibuprofen treatment, only the transductal diameter/kg was larger in the clinical symptoms group compared to the no clinical symptoms group. There was no difference in cardiomegaly or pulmonary edema on the chest radiograph between the two groups (Table 2). In the clinical symptoms group, 5 infants (35.7%) had both cardiomegaly and pulmonary edema on the chest radiograph; however, 7 infants (50.0%), had neither cardiomegaly nor pulmonary edema even though there were clinical symptoms due to hsPDA.

**Pharmacologic characteristics of ibuprofen treatment**

There was no significant difference in the median postnatal day of the first course of ibuprofen therapy in both groups. More infants received multiple courses of ibuprofen in the clinical symptoms group compared with the no clinical symptoms group (Table 3).

**Mortality and in-hospital outcomes**

PDA closure was confirmed in 86 (94.5%) infants in our study

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**Table 1. Demographic and baseline characteristics of the study population**

| Characteristics                        | hsPDA without clinical symptoms group (n = 77; 84.6%) | hsPDA with clinical symptoms group (n = 14; 15.4%) | \( P \) value |
|----------------------------------------|-----------------------------------------------------|---------------------------------------------------|---------------|
| GA, wk                                 | 26 [23–27]                                          | 25 [23–27]                                        | 0.012         |
| Birth weight                           | 880 [370–1,780]                                     | 655 [600–930]                                     | <0.001        |
| Male, gender                           | 39 (50.6%)                                          | 6 (42.9%)                                         | 0.592         |
| Year of birth ≥ 2013                   | 22 (28.6%)                                          | 9 (64.3%)                                         | 0.014         |
| Multiple gestation                     | 49 (63.6%)                                          | 7 (50.0%)                                         | 0.335         |
| Cesarean section                       | 37 (48.1%)                                          | 8 (57.1%)                                         | 0.531         |
| Histologic chorioamnionitis            | 43 (55.8%)                                          | 9 (64.3%)                                         | 0.557         |
| PH                                     | 6 (7.8%)                                            | 3 (21.4%)                                         | 0.139         |
| Antenatal steroid use, complete        | 44 (57.1%)                                          | 11 (78.6%)                                        | 0.131         |
| SGA                                    | 11 (14.3%)                                          | 5 (35.7%)                                         | 0.066         |
| DR resuscitation                      | 9 (11.7%)                                           | 4 (28.6%)                                         | 0.110         |
| 5-minute Apgar score                   | 6 (2–9)                                             | 5 (0–7)                                           | 0.235         |
| Surfactant use, any                    | 55 (71.4%)                                          | 13 (92.9%)                                        | 0.107         |
| Surfactant use, multiple               | 10 (13.0%)                                          | 2 (14.3%)                                         | 1.000         |
| CRIB-II score                          | 11 (8–16)                                           | 13 (9–18)                                         | 0.002         |
| Invasive ventilator care ≤ 2 days after birth | 54 (70.1%)                                    | 14 (100.0%)                                      | 0.017         |
| Hypotension before treatment           | 10 (13.0%)                                          | 11 (78.6%)                                        | <0.001        |
| INO use before treatment               | 4 (5.2%)                                            | 0 (0.0%)                                          | 1.000         |
| Sepsis before PDA treatment            | 2 (2.6%)                                            | 1 (7.1%)                                          | 0.398         |

\( hsPDA = \) hemodynamically-significant patent ductus arteriosus, GA = gestational age, PIH = pregnancy induced hypertension, SGA = small for gestational age, DR = delivery room, CRIB = clinical risk index for babies, INO = inhaled nitric oxide, PDA = patent ductus arteriosus.

**Table 2. Echocardiographic and radiographic characteristics of the study population**

| Characteristics                        | hsPDA without clinical symptoms group (n = 77; 84.6%) | hsPDA with clinical symptoms group (n = 14; 15.4%) | \( P \) value |
|----------------------------------------|-----------------------------------------------------|---------------------------------------------------|---------------|
| Transductal diameter per               |                                                    |                                                   |               |
| Weight, mm/kg                          | 2.8 [1.4–6.7]                                       | 3.9 [2.7–6.3]                                     | 0.001         |
| LA/Ao ratio, %                         | 1.6 [1.1–3.0]                                       | 1.7 [1.3–2.3]                                     | 0.157         |
| LVDd ratio, %                          | 111 [79–167]                                       | 111 [85–127]                                     | 0.183         |
| Radiographic characteristics           |                                                    |                                                   |               |
| Cardiomegaly                           | 29 (37.7%)                                          | 5 (35.7%)                                         | 0.890         |
| Pulmonary edema                        | 30 (39.0%)                                          | 7 (50.0%)                                         | 0.439         |

\( hsPDA = \) hemodynamically-significant patent ductus arteriosus, LA/Ao = left atrial/aortic root, LVDd = left ventricular end-diastolic dimension.
Thirty-four infants (37.4%) received secondary PDA ligation without clinical symptoms group. Six (7.8%) infants received secondary PDA ligation without clinical symptoms group. Multivariate analysis of mortality and in-hospital outcomes according to the presence of clinical symptoms due to PDA at the time of ibuprofen treatment (reference: hsPDA without clinical symptoms group) indicated that secondary PDA ligation after failure of PDA closure by ibuprofen use was higher in the clinical symptoms group compared with the no clinical symptoms group in the univariate analysis (Table 4).

Three infants died after PDA closure confirmed by echocardiography. One infant died by multi-organ failure due to NEC 3b, and the other 2 infants died because of sepsis from methicillin-resistant *Staphylococcus aureus* and imipenem-resistant *Acinetobacter baumannii* in each case, respectively. In the univariate analysis, there were no significant differences in the mortality; however, the total duration of hospitalization, invasive ventilator care, and total respiratory support were all longer, and BPD was more frequent in the clinical symptoms group compared with the no clinical symptoms group in the univariate analysis (Table 4).

After multivariate analysis adjusting for the CRIB-II score, birthweight, the year of birth, and the invasive ventilator care ≤ 2 postnatal days, there were no significant differences in mortality, secondary PDA ligation and in-hospital outcomes between the two groups (13 [3–89] days in the no clinical symptoms group vs. 18 [3–41] days in the clinical symptoms group; P = 0.837). Thirty-four infants (37.4%) received secondary PDA ligation after failure of PDA closure by ibuprofen use. The frequency of secondary PDA ligation was higher in the clinical symptoms group compared with the no clinical symptoms group in the univariate analysis.

Table 3. Pharmacological characteristics of ibuprofen treatment in the study population

| Characteristics                        | hsPDA without clinical symptoms group (n = 77; 84.6%) | hsPDA with clinical symptoms group (n = 14; 15.4%) | P value |
|----------------------------------------|------------------------------------------------------|--------------------------------------------------|---------|
| Postnatal day of first day of ibuprofen treatment | 2 (0–21)                                              | 3 (1–11)                                         | 0.340   |
| Route of ibuprofen administration      |                                                      |                                                  |         |
| Intravenous ibuprofen                  | 57 (74.0%)                                            | 11 (78.6%)                                       | 1.000   |
| Oral ibuprofen                         | 20 (26.0%)                                            | 3 (21.4%)                                        |         |
| No. of courses                         |                                                      |                                                  | 0.047   |
| 1                                      | 53 (68.8%)                                            | 5 (35.7%)                                        |         |
| 2                                      | 18 (23.4%)                                            | 7 (50.0%)                                        |         |
| 3                                      | 6 (7.8%)                                              | 2 (14.3%)                                        |         |

hsPDA = hemodynamically-significant patent ductus arteriosus.

Table 4. In-hospital outcomes according to the presence of clinical symptoms due to PDA at the time of ibuprofen treatment

| Outcomes                                            | hsPDA without clinical symptoms group (n = 77; 84.6%) | hsPDA with clinical symptoms group (n = 14; 15.4%) | P value |
|-----------------------------------------------------|------------------------------------------------------|--------------------------------------------------|---------|
| Mortality ≥ 3 days after birth                       | 2 (2.6%)                                             | 1 (7.1%)                                         | 0.398   |
| Confirmed PDA closure                                | 74 (96.1%)                                           | 12 (85.7%)                                       | 0.168   |
| Secondary PDA ligation                              | 25 (32.5%)                                           | 9 (64.3%)                                        | 0.024   |
| NEC after PDA treatment                              | 8 (10.4%)                                            | 2 (14.3%)                                        | 0.649   |
| SIP after PDA treatment                              | 2 (2.6%)                                             | 1 (7.1%)                                         | 0.398   |
| Sepsis after PDA treatment                          | 24 (31.2%)                                           | 4 (28.6%)                                        | 1.000   |
| IVH after PDA treatment                              | 6 (7.8%)                                             | 0 (0.0%)                                         | 0.585   |
| PVL after PDA treatment                              | 6 (7.8%)                                             | 1 (7.1%)                                         | 1.000   |
| Moderate to severe BPD                               | 41 (64.7%)                                           | 11 (84.6%)                                       | 0.043   |
| Severe BPD                                           | 16 (21.3%)                                           | 7 (53.8%)                                        | 0.034   |
| ROP requiring surgery or VEGF                        | 28 (37.3%)                                           | 6 (46.2%)                                        | 0.547   |
| Hospital days                                        | 87 (16–197)                                          | 107 (11–151)                                     | 0.043   |
| Days of invasive ventilator care                    | 20 (0–137)                                           | 56 (11–148)                                      | 0.001   |
| Days of respiratory support                         | 69 (16–189)                                          | 102 (11–151)                                     | 0.015   |
| Discharge with oxygen or home ventilator            | 23 (29.9%)                                           | 9 (64.3%)                                        | 0.030   |

PDA = patent ductus arteriosus, hsPDA = hemodynamically-significant PDA, NEC = necrotizing enterocolitis, SIP = spontaneous intestinal perforation, IVH = intraventricular hemorrhage, PVL = periventricular leukomalacia, BPD = bronchopulmonary dysplasia, ROP = retinopathy of prematurity, VEGF = vascular endothelial growth factor.

Table 5. Multivariate analysis of mortality and in-hospital outcomes according to the presence of clinical symptoms at the time of ibuprofen treatment (reference: hsPDA without clinical symptoms group)

| Variables                                             | hsPDA with clinical symptoms group | Adjusted OR | CI          | P value |
|-------------------------------------------------------|-----------------------------------|-------------|-------------|---------|
| Primary outcome                                       |                                    |             |             |         |
| Mortality ≥ 3 days after birth                         | 0.50                              | 0.005–52.352| 0.771       |         |
| Composite outcomea                                     | 1.10                              | 0.189–6.256 | 0.918       |         |
| Secondary PDA ligation                                | 0.85                              | 0.203–3.533 | 0.821       |         |
| Secondary outcome                                     | 3.40                              | 0.874–13.208| 0.077       |         |

Adjusted for the CRIB-II score, the year of birth, birthweight, invasive ventilator care ≤ 2 days after birth.

hsPDA = hemodynamically-significant patent ductus arteriosus, OR = odds ratio, CI = confidence interval, PDA = patent ductus arteriosus, NEC = necrotizing enterocolitis, IVH = intraventricular hemorrhage, BPD = bronchopulmonary dysplasia, *Mortality ≥ 3 days after birth, NEC, IVH, BPD; *Mortality ≥ 3 days after birth, NEC, IVH, severe BPD.

Population. There were no statistically significant differences in the PDA closure rate and the postnatal day of PDA closure between the two groups (13 [3–89] days in the no clinical symptoms group vs. 18 [3–41] days in the clinical symptoms group; P = 0.837). Thirty-four infants (37.4%) received secondary PDA ligation after failure of PDA closure by ibuprofen use. The frequency of secondary PDA ligation was higher in the clinical symptoms group compared with the no clinical symptoms group in the univariate analysis (Table 4).
the clinical symptoms group and the no clinical symptoms group (Table 5).

DISCUSSION

In our study, infants who received first ibuprofen treatment when there were clinical symptoms attributable to hsPDA were younger with higher CRIB-II scores than infants with first ibuprofen treatment done with no clinical symptoms due to hsPDA. More infants received multiple courses of ibuprofen treatment who received first ibuprofen treatment when there were clinical symptoms attributable to hsPDA; however, there were no significant differences in the mortality and in-hospital outcomes between the two groups.

All the patent ductus are not pathologic. In some cases, the patent ductus may be an “innocent physiologic bystander” with little hemodynamic consequences; however, in other cases, it may contribute to significant pathologic changes for which early detection and intervention are warranted to prevent neonatal morbidity (4,20). We usually define hsPDA as PDA with a left-to-right shunt from the aorta into the pulmonary arteries which causes pulmonary vascular and left ventricular volume overload. However, there is no consensus on the definition of hsPDA (9,21). The lack of a standardized approach in determining the hemodynamic significance is a major barrier in searching for an association between persistent ductus arteriosus and the major morbidities of preterm infants or in assessing the benefit of PDA treatment (8). Before the PDA staging system was proposed by McNamara and Seghal (12), a transductal diameter more than 1.5 mm was proposed as a significant factor implying end-organ hypoperfusion reported by Kluckow and Evans (11). McNamara and Seghal (12,22) proposed the staging of PDA for determining the severity of hsPDA, which was based on the clinical and echocardiographic criteria. Echocardiographic staging includes E1 (no evidence of ductal flow), E2 (small non-significant ductus arteriosus), E3 (moderate hsPDA), and E4 (large hsPDA). Clinical staging includes C1 (asymptomatic), C2 (mild), C3 (moderate), and C4 (severe) based on illness severity and the magnitude of the cardiovascular, respiratory, and gastrointestinal problems.

Our unit adapted the definitions for hsPDA according to the suggestions of Kluckow and Evans (11) and McNamara and Seghal (12,22) if PDA with a transductal diameter ≥ 1.4 mm/kg with a significant left to right shunt was confirmed by the echocardiography. Because our study was a retrospective study, we could not get the whole echocardiographic parameters such as ductal flow pattern, early passive to late atrial contractile phase of transmitial filling ratio, and isovolumic relaxation time. Thus, we could not define the hsPDA entirely dependent on the McNamara staging based on echocardiographic criteria. For the definition of clinical symptoms due to hsPDA, our definition of clinical symptoms was close to C3 or C4 of clinical staging by McNamara and Seghal (12,22). When we tried to classify our study population according to the McNamara’s clinical staging, it was difficult to judge which group is suitable for each baby because some parameter fitted one stage, however, other parameters fitted another. Especially about C2 and C3 group, most infants were in the gray zone between C2 and C3 group, so we decided to adapt another guideline to define clinical symptoms due to PDA according to the Sosenko et al. (15). We did not include infants with mild respiratory symptoms in the clinical symptoms group because mild respiratory symptoms can be due to other causes such as respiratory distress syndrome, sepsis or “evolving” BPD.

There have been many reports suggesting a correlation between the increase of the plasma BNP level and the echocardiographic parameter suggesting hsPDA or symptomatic PDA (10,23). Choi et al. (24) suggested that the BNP levels correlated well with the magnitude of the ductal shunt, and the BNP levels were higher in the symptomatic PDA group when compared with the asymptomatic PDA group. Other researchers suggested that the early BNP level was significantly correlated with the magnitude of the ductal shunt and can help to predict hsPDA (25,26). Because we could get the BNP level in only 14 infants among the study population (data are not shown), we could not examine such a correlation between the BNP level and other echocardiographic or radiographic parameters or the association between the severity of the clinical symptoms and the BNP level. Further studies are needed focusing on the association between the BNP level and hsPDA.

In our study, only the transductal diameter was a significant echocardiographic parameter which can predict the presence of clinical symptoms due to hsPDA. However, when we assessed the correlation between the findings of the radiography and the clinical symptoms, there was no significant association in our study. Because the size of the thymus of a neonate is relatively big and chest radiography is usually done as an anteroposterior view, we could not obtain exact information about cardiomegaly, and echocardiographic measurement can be a better method to judge cardiomegaly. There are several studies on significant echocardiographic parameters as an early marker of hsPDA because echocardiographic signs usually have been shown to precede clinical symptoms by 1.8 days (27). Some researchers have suggested that a transductal diameter exceeding 1.5 mm combined with diastolic flow reversal in the descending aorta may be the best echocardiographic criterion (28). Others have stated that a substantial ductal shunt is associated with the following: an increased ratio of left atrial to aortic root dimensions ≥ 1.5:1, a ductal diameter ≥ 1.5 mm, a measured IVDD ratio by the normal LVDD ratio of more than 115% (29), left ventricular volume and pressure overloading, and reversal of diastolic flow in the descending aorta or in the cerebral or re-
nal arteries (12,30). About left ventricular volume and pressure overloading, an increased LVDD ratio (measured LVDD divided by the normal LVDD as an index) > 115% was a useful index of PDA treatment (29). However, in isolation, many of these markers have low sensitivity and specificity for ductal significance when compared to the transdural diameter (11). Therefore, a study on the combination of clinical and echocardiographic markers is needed for better decision making about PDA treatment.

There still is no consensus on which PDAs we have to treat, when to treat, and how to treat. There is a high chance of spontaneous closure of preterm PDA, especially in infants with higher GA (21,28). Moreover, there is some evidence that preterm infants with mild symptoms from PDA do not necessarily benefit from early pharmacologic treatment compared with conservative treatment (15,31). Additionally, there are potential risks associated with pharmacologic treatment, such as impaired renal function, intestinal perforation, NEC and increase of bleeding tendency. Many studies on surgical ligation have reported adverse outcomes such as increased BPD in association with surgical ligation (32). The timing of PDA ligation is also important for determining the neonatal outcomes (33). Thus, targeted treatment is important to reduce such adverse effects, and knowing early markers of hsPDA that indicate the need for treatment will be beneficial.

In our study population, the infants in the clinical symptoms group were younger and the CRIB-II score was higher; and after adjusting for such differences in the baseline characteristics, there were no significant differences in the mortality and in-hospital outcomes between the two groups. Although more infants received repeated courses of ibuprofen treatment in the clinical symptoms group, there were no significant differences in the PDA closure rate and in the duration of PDA opening. Previous studies have suggested that ductal ligation causes an abrupt change in the hemodynamics of preterm infants, thereby leading to adverse neonatal outcomes (34). Recently, Lemmers et al. (35) performed near infrared spectroscopy-monitored cerebral oxygen saturation (rSco2) and brain MRI at term equivalent age and assessed the regional brain volume and maturation of the posterior limb of the internal capsule in an indomethacin group, PDA ligation group, and matched control groups without PDA. PDA led to lower cerebral oxygenation, and the PDA ligation group had the lowest rSco2 values, the highest postnatal age before effective treatment, and the lowest volumes of most brain regions, especially the cerebellum. We can infer that the longer period of suboptimal cerebral oxygenation in the PDA ligation group, not the surgical procedure itself, could be a reason for the negative effect of PDA on brain growth. We can also guess that current practice to postpone medical or surgical ductal closure may cause negative effects on brain growth. In our study, the frequency of secondary PDA ligation was somewhat higher in the clinical symptoms group with marginal statistical significance after multivariate analysis; however, it did not affect the mortality and the frequency of BPD. And there was no difference in the postnatal day of PDA closure between two groups. Our data favor the selected treatment of clinically symptomatic PDA without adverse clinical outcomes.

In our study, there were no significant differences in mortality or neonatal outcomes between prophylactic or pre-symptomatic treatment and symptomatic treatment. Although the meta-analysis suggests that prophylactic indomethacin treatment can reduce the frequency of IVH and reduce later PDA treatment and PDA ligation, there was no significant difference in mortality or long-term neurodevelopmental outcomes (36). Yeo et al. (37) reported that prophylactic ibuprofen treatment in preterm infants could not reduce PDA ligation and the incidence of IVH. Concerning pre-symptomatic treatment, there are some reports which compared the mortality and neonatal outcomes between early and conventional treatment groups. Van Overmeire et al. (38) reported that early treatment at postnatal day 3 was associated with more renal side effects without any advantage in respiratory or major outcomes. Candel-Pau et al. (39) performed routine echocardiography within 48 hours after birth in infants < 29 weeks of gestation, and if the PDA diameter/weight ≥ 1.4 mm/kg and/or the LA/Ao ratio ≥ 1.4, indomethacin treatment was done. However, when they compared the frequency of secondary ligation, mortality and major preterm morbidities between such an early treatment group and the conventional treatment group, there were no significant differences like our study results.

Our study has some limitations. First, this is a retrospective study with a small number of patients in a single center. Thus, the factors with marginal statistical significance like secondary PDA treatment should be reassessed with a large cohort. Actually, ibuprofen was recently accepted as a treatment modality of preterm PDA and we included all the infants with ibuprofen treatment after intravenous type of ibuprofen became available in Korea. So, although the power level of our study did not reach 0.80, the results of our study are meaningful as a suggestion on the needs of future prospective, multi-center study. Second, we only included the preterm infants with ibuprofen treatment and the treatment policy changed from 2013. Moreover, we could not assess the outcome of infants with conservative management for preterm PDA which is an increasing treatment policy for preterm PDA after the year 2013. Although we tried to adjust such bias by year factor in multiple regression analysis in this study, further studies are needed focusing on the association between conservative treatment of preterm PDA and neonatal outcomes. Third, the definition of clinical symptoms from hsPDA was somewhat different from the clinical staging of preterm PDA by McNamara and Sehgal (12,22). If we could classify the group according to their clinical staging, we could assess the relationship between the severity of the clinical symptoms and
the neonatal outcomes in preterm infants, so further studies are needed. In our study, about 15% of the infants with ibuprofen treatment received treatment in situations for which the clinical symptoms were attributable to the hsPDA. Younger infants tended to receive first ibuprofen treatment when there were clinical symptoms due to hsPDA. Although more infants received multiple courses of ibuprofen treatment in the clinical symptoms group, there were no differences in the mortality and in-hospital outcomes between the infants with clinical symptoms due to hsPDA and the infants without clinical symptoms. We can hold off on PDA treatment until the clinical symptoms becomes prominent. However, because our study was a retrospective, single center study with small number of study objects, a prospective, multicenter, large cohort study is needed in the near future.

DISCLOSURE

The authors have no potential conflicts of interest to disclose.

AUTHOR CONTRIBUTION

Conception and Design of the study: Lee JA, Jung YH, Kim HS. Acquisition of data: Yoo H. Statistical analysis: Oh S. First draft of the manuscript: Yoo H, Lee JA. Revision and critical review of the manuscript: Yoo H, Lee JA, Jung YH, Sohn JA, Shin SH, Choi CW, Kim EK, Kim HS, Kim BI. Manuscript approval: all authors.

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REFERENCES

1. Nemerofsky SL, Parravicini E, Bateman D, Kleinman C, Polin RA, Lorenz JM. The ductus arteriosus rarely requires treatment in infants > 1000 grams. Am J Perinatol 2008; 25: 661-6.
2. Clyman RI, Couto J, Murphy GM. Patent ductus arteriosus: are current neonatal treatment options better or worse than no treatment at all? Semin Perinatol 2012; 36: 123-9.
3. Heuchan AM, Clyman RI. Managing the patent ductus arteriosus: current treatment options. Arch Dis Child Fetal Neonatal Ed 2014; 99: F431-6.
4. Evans N. Preterm patent ductus arteriosus: a continuing conundrum for the neonatologist? Semin Fetal Neonatal Med 2015; 20: 272-7.
5. Clyman RI. The role of patent ductus arteriosus and its treatments in the development of bronchopulmonary dysplasia. Semin Perinatol 2013; 37: 102-7.
6. Benitz WE. Treatment of persistent patent ductus arteriosus in preterm infants: time to accept the null hypothesis? J Perinatol 2010; 30: 241-52.
7. Lee JA, Kim MJ, Oh S, Choi BM. Current status of therapeutic strategies for patent ductus arteriosus in very-low-birth-weight infants in Korea. J Korean Med Sci 2015; 30 Suppl 1: S59-66.
8. Benitz WE; Committee on Fetus and Newborn, American Academy of Pediatrics. Patent ductus arteriosus in preterm infants. Pediatrics 2016; 137: 1-6.
9. Zonnenberg I, de Waal K. The definition of a haemodynamic significant duct in randomized controlled trials: a systematic literature review. Acta Paediatr 2012; 101: 247-51.
10. Flynn PA, da Graca RL, Auld PA, Nesin M, Kleinman CS. The use of a bed-side assay for plasma B-type natriuretic peptide as a biomarker in the management of patent ductus arteriosus in premature neonates. J Pediatr 2005; 147: 38-42.
11. Kluckow M, Evans N. Early echocardiographic prediction of symptomatic patent ductus arteriosus in preterm infants undergoing mechanical ventilation. J Pediatr 1995; 127: 774-9.
12. McNamara PJ, Sehgal A. Towards rational management of the patent ductus arteriosus: the need for disease staging. Arch Dis Child Fetal Neonatal Ed 2007; 92: F424-7.
13. Schena F, Francescato G, Cappelleri A, Picciolli I, Mayer A, Mosca E, Fumagalli M. Association between hemodynamically significant patent ductus arteriosus and bronchopulmonary dysplasia. J Pediatr 2015; 166: 1488-92.
14. Parry G, Tucker J, Tarnow-Mordi W; UK Neonatal Staffing Study Collaborative Group. CRIB II: an update of the clinical risk index for babies score. Lancet 2003; 361: 1789-91.
15. Sosenko IR, Fajardo MF, Claire N, Bancalari E. Timing of patent ductus arteriosus treatment and respiratory outcome in premature infants: a double-blind randomized controlled trial. J Pediatr 2012; 160: 929-935. e1.
16. Olsen IE, Grovenman SA, Lawson ML, Clark RH, Zemel BS. New intrauterine growth curves based on United States data. Pediatrics 2010; 125: e214-24.
17. Bell MJ, Ternberg JL, Feigin RD, Keating JP, Marshall R, Barton L, Brotheron T. Neonatal necrotizing enterocolitis. Therapeutic decisions based upon clinical staging. Ann Surg 1978; 187: 1-7.
18. Jobe AH, Bancalari E. Bronchopulmonary dysplasia. Am J Respir Crit Care Med 2001; 163: 1723-9.
19. Papile LA, Burstein J, Burstein R, Koffler H. Incidence and evolution of subependymal and intraventricular hemorrhage: a study of infants with birth weights less than 1,500 gm. J Pediatr 1978; 92: 529-34.
20. Bose CL, Laughon MM. Patent ductus arteriosus: lack of evidence for common treatments. Arch Dis Child Fetal Neonatal Ed 2007; 92: F498-502.
21. Sullivan H, Koehne P, Hansmann G. Recent advances in the treatment of preterm newborn infants with patent ductus arteriosus. Clin Perinatol 2016; 43: 113-29.
22. Sehgal A, McNamara PJ. Does echocardiography facilitate determination...
of hemodynamic significance attributable to the ductus arteriosus? Eur J Pediatr 2009; 168: 907-14.
23. Jeong HA, Shin J, Kim E, Lee EH, Choi BM, Son CS, Lee JW. Correlation of B-type natriuretic peptide levels and echocardiographic parameters in preterm infants with patent ductus arteriosus. Korean J Pediatr 2016; 59: 183-9.
24. Choi BM, Lee KH, Eun BL, Yoo KH, Hong YS, Son CS, Lee JW. Utility of rapid B-type natriuretic peptide assay for diagnosis of symptomatic patent ductus arteriosus in preterm infants. Pediatrics 2005; 115: e255-61.
25. Lee JH, Shin JH, Park KH, Rihe YJ, Park MS, Choi BM. Can early B-type natriuretic peptide assays predict symptomatic patent ductus arteriosus in extremely low birth weight infants? Neonatology 2013; 103: 118-22.
26. Kim JS, Shim EJ. B-type natriuretic Peptide assay for the diagnosis and prognosis of patent ductus arteriosus in preterm infants. Korean Circ J 2012; 42: 192-6.
27. Skelton R, Evans N, Smythe J. A blinded comparison of clinical and echocardiographic evaluation of the preterm infant for patent ductus arteriosus. J Paediatr Child Health 1994; 30: 406-11.
28. Jain A, Shah PS. Diagnosis, evaluation, and management of patent ductus arteriosus in preterm neonates. JAMA Pediatr 2015; 169: 863-72.
29. Nagasawa H, Terazawa D, Kohno Y, Yamamoto Y, Kondo M, Sugawara M, Koyama T, Miura R. Novel treatment criteria for persistent ductus arteriosus in neonates. Pediatr Neonatol 2014; 55: 250-5.
30. El Hajjar M, Vaksmann G, Rakza T, Kongolo G, Storme L. Severity of the ductal shunt: a comparison of different markers. Arch Dis Child Fetal Neonatal Ed 2005; 90: F419-22.
31. Vanhaesebrouck S, Zonnenberg I, Vandervoort P, Brunee E, Van Hoestenberghe MR, Theyskens C. Conservative treatment for patent ductus arteriosus in the preterm. Arch Dis Child Fetal Neonatal Ed 2007; 92: F244-7.
32. Mirea L, Sankaran K, Seshia M, Ohlsson A, Allen AC, Aziz K, Lee SK, Shah PS; Canadian Neonatal Network. Treatment of patent ductus arteriosus and neonatal mortality/morbidities: adjustment for treatment selection bias. J Pediatr 2012; 161: 689-694.e1.
33. Sung SI, Choi SY, Park JH, Lee MS, Yoo HS, Ahn SY, Chang YS, Park WS. The timing of surgical ligation for patent ductus arteriosus is associated with neonatal morbidity in extremely preterm infants born at 23-25 weeks of gestation. J Korean Med Sci 2014; 29: 581-6.
34. Kabra NS, Schmidt B, Roberts RS, Doyle LW, Papile L, Fanaroff A; Trial of Indomethacin Prophylaxis in Preterms Investigators. Neurosensory impairment after surgical closure of patent ductus arteriosus in extremely low birth weight infants: results from the trial of indomethacin prophylaxis in preterms. J Pediatr 2007; 150: 229-34, 234.e1.
35. Lemmers PM, Benders MJ, D’Ascenzo R, Zethof J, Alderliesten T, Kersbergen KJ, Issum I, de Vries LS, Groenendaal F, van Bel F. Patent ductus arteriosus and brain volume. Pediatrics 2016; 137; e20153090.
36. Fowlie PW, Davis PG, McGuire W. Prophylactic intravenous indomethacin for preventing mortality and morbidity in preterm infants. Cochrane Database Syst Rev 2010: CD000174.
37. Yeo MS, Choi K, Lee HJ, Park HK, Kim CR, Seol II. Effect of prophylactic ibuprofen in preterm infants less than 1,250 g in birth weight. J Korean Soc Neonatol 2011; 18: 234-9.
38. Van Overmeire B, Van de Broek H, Van Laer P, Weyler J, Vanhaesebrouck P. Early versus late indomethacin treatment for patent ductus arteriosus in premature infants with respiratory distress syndrome. J Pediatr 2001; 138: 205-11.
39. Candel-Pau J, Linde Sillo Á, Castillo Salinas F, Coma Redon E, Ferrer Menduña Q, Albert DC. Early versus conventional treatment for patent ductus arteriosus in preterm infants. Am J Perinatol 2013; 30: 289-95.