Can red cell distribution width in very low birth weight infants predict bronchopulmonary dysplasia?

Seong Hee Oh, MDa, Hyun-Jeong Do, MDa, Ji Sook Park, MD, PhDb,c, Jae Young Cho, MDb,c, Chan-Hoo Park, MD, PhDc,d,*

Abstract
Red cell distribution width (RDW) is a useful marker for assessing the severity and prognosis of various diseases in adults. However, whether it is applicable to children, especially in newborns, has not been determined. This study aimed to investigate the RDW values of preterm infants and evaluate whether RDW values in the early days of life can predict bronchopulmonary dysplasia (BPD) development.

One hundred and eight infants born at <30 weeks of gestation with a birth weight of <1500 g participated in this retrospective study. RDW values measured at birth, 7 days (D7), and 28 days (D28) after birth were reviewed. The changes in RDW values in the first month of life were analyzed, and we evaluated the relationship between RDW and BPD.

The mean RDW values at birth, D7, and D28 and the change from birth to D7 were 16.2 ± 0.1%, 17.5 ± 0.2%, 17.6 ± 0.2% and 1.3 ± 1.8%, respectively. RDW at birth was lower in the infants born at <28 weeks’ gestational age than in those born at ≥28 weeks’ gestational age (15.7 ± 0.3 vs 16.4 ± 0.2, P = .024). RDW values of both groups increased during the first week after birth and did not differ significantly at D7. The levels remained similar at 1 month of age. RDW at birth, D7, and D28 and the changes in RDW from birth to D7 were not correlated with the development of BPD independent of its severity.

The usefulness of RDW as a predictor of BPD development remains questionable and requires further study.

Abbreviations: BPD = bronchopulmonary dysplasia, CBC = complete blood cell count, D28 = twenty-eighth day of life, D7 = seventh day of life, GA = gestational age, RBC = red blood cell, RDW = red cell distribution width.

Keywords: bronchopulmonary, dysplasia, infant, premature, red cell distribution width, very low birth weight infant

1. Introduction

Red cell distribution width (RDW) as part of complete blood count (CBC) has traditionally been used with mean corpuscular volume (MCV) to determine the cause of anemia. RDW values increase in circumstances of ineffective erythropoiesis in the bone marrow such as iron deficiency anemia, folate and vitamin B12 deficiency, or shortening of the red blood cell (RBC) life span by destruction such as in sickle cell anemia.[1]

Many recent studies have reported that RDW is related to inflammation[2–4] and hypoxemia,[5] and is an easily available parameter that can predict the severity[6,7] and prognosis of various diseases in adults.[15,17] Additional blood sampling is not required to determine the RDW values, as CBC is performed relatively frequently and requires only a small amount of blood. Therefore, RDW might be a useful tool for assessing the medical conditions of newborns, especially preterm infants. However, there are no sufficient studies in children and infants on this topic.[18–23] Moreover, there are few studies on preterm infants, and the normal range of RDW values in preterm infants has yet to be determined.[24–26]

Bronchopulmonary dysplasia (BPD), which is a major complication of preterm infants, occurs when premature lung tissue and vessels are injured and their development and differentiation are disrupted by perinatal inflammation, such as chorioamnionitis, hyperoxia, mechanical ventilation, and infection. There have been a few previous studies on the
relationship between RDW and BPD, but they have shown conflicting results. The authors investigated the changes in RDW values in very low birth weight infants born before 30 weeks of gestation and whether the RDW values measured in the early days of life can predict BPD development.

2. Methods

This study was a retrospective review of the medical records of patients hospitalized in the two-level III neonatal intensive care units at Gyeongsang National University Hospital and Changwon Gyeongsang National University Hospital in South Korea between January 2009 and August 2019. This study was approved by the Institutional Review Boards of the Gyeongsang National University Hospital (IRB no. 2020-02-014) and Changwon Gyeongsang National University Hospital (IRB no. 2020-02-017), which waived the need for informed consent. All methods were performed in accordance with the relevant guidelines and regulations.

Patients born at gestational age (GA) <30 weeks and birth weight <1500 g were included in this study. Patients with 1. a chromosomal abnormality or major congenital anomaly; 2. culture-proven early-onset sepsis; 3. a recent RBC transfusion (within 1 week after birth); or 4. maternal anemia (hemoglobin <8 g/dL) were excluded.

RDW values measured at birth, 7 days (D7), and 28 days (D28) after birth were reviewed. The RDW levels according to GA and the association between RDW and BPD development were analyzed. RDW was determined as part of a CBC, which was checked within 1 hour after birth, at D7, D28, and per the local routine protocol as necessary. The 0.3 to 0.5 mL of Blood samples were taken from the radial or umbilical artery and placed in ethylenediaminetetraacetic acid-containing tubes. CBC was measured using an automated hematology analyzer (Sysmex, Kobe, Japan) and quality controls and calibrations were performed regularly according to standard rules. White blood cell counts, hemoglobin, MCV, and platelet counts were also collected using the RDW from the CBC.

BDP was defined according to the National Institute of Child Health and Human Development consensus definition. BPD was diagnosed if the patient required artificial respiratory support providing positive airway pressure or oxygen for more than 28 days. Moderate or severe BPD was diagnosed if the patient needed positive airway pressure or oxygen at 36 weeks’ postmenstrual age or at discharge. Babies who died in the first month of life were excluded from the assessment of the relationship between RDW values and BPD development.

Statistical analysis was performed using SAS software (version 9.4; SAS Institute, Inc., Cary, NC, USA). The normality of continuous values was assessed using the Kolmogorov-Smirnov test. Categorical and continuous variables are presented as number (percentage) and mean ± standard deviation or median (interquartile range [IQR]), respectively. Categorical variables were analyzed using the Chi-Squared test or Fisher exact test. Continuous variables were analyzed using the Student t test or Mann–Whitney’s U test. A linear mixed-effects model was used to analyze the trends of RDW during the first month of life and compared them between infants born at <28 weeks’ GA and those born at ≥28 weeks’ GA. Multivariable logistic regression was used to analyze the association between RDW and BPD. Statistical significance was set at P < .05.

3. Results

3.1. Characteristics of the infants

Of the 171 infants eligible for this study, 63 were excluded. The mean GA and birth weight of the remaining 108 patients were 28.4 ± 1.4 weeks’ gestation and 1165.6 ± 212.9 g. The number of infants with BPD and moderate/severe BPD were 64 (59.3%) and 21 (19.4%), respectively. White blood cell count and C-reactive protein were higher in infants born at <28 weeks’ GA (n = 27) than 28 to 29 weeks’ GA (n = 81). BPD and moderate/severe BPD presented more in the infants born before 28 weeks of gestation than in the infants born at 28 weeks of gestation or later (Table 1).

3.2. RDW values in preterm infants

The mean RDW values at birth, D7, and D28 are presented in Table 2. The mean RDW values at birth, D7, and D28 were 16.2 ± 6.4, 15.4 ± 6.8, and 15.6 ± 8.9, respectively. The RDW value at birth was lower in infants born at <28 weeks’ GA than in infants born at ≥28 weeks’ GA.

Table 1
Demographic characteristics and laboratory data by gestational age.

| GA < 28 weeks (n=27) | GA 28–29 weeks (n=81) | P value |
|----------------------|------------------------|--------|
| Birth weight (g)     | 976.9±213.2            | 1228.5±172.8 | <.001 |
| SGA (n, %)           | 3 (11.1)               | 9 (11.1)     | 1.000 |
| Male (n, %)          | 14 (51.9)              | 54 (66.7)    | .167 |
| C-sec (n, %)         | 16 (59.3)              | 62 (76.5)    | .135 |
| Maternal hypertension (n, %) | 5 (18.5) | 18 (22.2) | .684 |
| WBC (10³/µL)         | 11.1 (5.9–15.9)        | 5.7 (4.1–8.2) | <.001 |
| Hemoglobin (g/dL)    | 15.5±1.0               | 16.0±1.6     | .059 |
| MCV (fl)             | 115.6±6.4              | 115.4±6.0    | .955 |
| Platelet (10³/µL)    | 256.1±69.1             | 232.2±67.8   | .118 |
| CRP (mg/L)           | 0.3 (0.2–0.4)          | 0.2 (0.1–0.2) | .003 |
| BPD (n, %)           | 21 (91.3)              | 43 (53.1)    | <.001 |
| Moderate BPD (n, %)  | 12 (52.2)              | 9 (11.1)     | <.001 |

BDP = bronchopulmonary dysplasia, CRP = C-reactive protein, C-sec = cesarean section, MCV = mean corpuscular volume, SGA = small for gestational age, WBC = white blood cell count.

Data are presented as mean ± standard deviation or median (interquartile range), and number (%).

* The data of 4 patients expired within 28 days of life were missing.

Oh et al. Medicine (2022) 101:3
than in those born at ≥28 weeks’ GA (P=.024). The RDW of both groups increased in the first week of life (P<.001, both) and then remained at similar levels for a month after birth (P=.639 and P=.772, respectively) without differences between the groups (Table 2, Fig. 1).

### 3.3. Relationship between RDW and BPD

After 4 patients who died in the first month of life were excluded, the relationship between RDW and BPD development in the remaining 104 patients was evaluated. The RDW values at birth, D7, the change of RDW between birth and D7, and D28 did not show any difference in accordance to BPD and moderate/severe BPD. These results remained similar even after we divided the patients into 2 groups born <28 weeks of GA and 28 to 29 weeks of GA. (Table 3). In multivariable logistic regression, the RDW values were not related to BPD presentation regardless of BPD severity (Table 4).

### 4. Discussion

The present study aimed to determine the changes in RDW values in preterm infants born at <30 weeks’ gestation and the relationship between RDW and BPD. The RDW values of infants born at <28 weeks’ GA were lower than those of infants born at 28 to 29 weeks’ GA. The RDW values in both groups increased during the first week of life, reached similar values, and remained similar throughout the first month of life. The RDW values in a month of life did not differ significantly between BPD groups.

The RDW levels of newborns are higher than those of children or adults due to active erythropoiesis and physiologic reticulocytosis.[29,30] However, it is challenging to determine the normal range of RDW in preterm infants because they are changing physiologically during the perinatal period and are influenced by various conditions such as GA. The upper reference was higher in

### Table 2

|                          | At birth | D7 | D28 | P value |
|--------------------------|----------|----|-----|---------|
| **Interaction effect**   |          |    |     |         |
| Overall                  | <0.001   | <0.001 | 0.658 |
| At birth vs D7           | <0.001   | <0.001 | 0.658 |
| D7 vs D28                | <0.001   | <0.001 | 0.658 |
| **Group effect**         |          |    |     |         |
| All                      | 0.024    | 0.832 | 0.501 |
| GA <28                   | 0.024    | 0.832 | 0.501 |
| GA ≥28                   | 0.024    | 0.832 | 0.501 |

D28 = twenty-eighth day of life, D7 = seventh day of life, GA = gestational age, RDW = red cell distribution width.

Data are presented as mean ± standard deviation.

### Table 3

|                     | Total | GA 28 weeks | GA 28–29 weeks |
|---------------------|-------|-------------|---------------|
|                     | BPD (n=64) | NonBPD (n=40) | P value | BPD (n=21) | NonBPD (n=2) | P value | BPD (n=43) | NonBPD (n=38) | P value |
| RDW at birth        | 16.0 (15.5–16.8) | 15.9 (15.3–16.7) | .390 | 15.8 (15.3–16.3) | 15.9 | .783 | 16.3 (15.5–17.0) | 15.9 (15.2–16.7) | .138 |
| RDW at D7           | 17.3 (16.1–18.4) | 16.8 (15.7–17.6) | .167 | 17.0 (16.0–18.9) | 18.5 | .561 | 17.3 (16.2–18.3) | 16.8 (15.6–17.6) | .105 |
| RDW at D28          | 17.8 (16.7–18.9) | 17.1 (16.2–18.7) | .282 | 17.9 (17.2–17.9) | 17.4 | .640 | 17.8 (16.6–18.8) | 17.1 (16.2–18.8) | .382 |
| RDW birth-D7        | 0.9 (0.3–2.1) | 0.8 (0.1–1.2) | .299 | 1.0 (0.3–2.7) | 2.6 | .522 | 0.8 (0.3–1.8) | 0.8 (0.1–1.1) | .349 |

|                     | Moderate/ severe BPD (n=21) | Non- moderate/ severe BPD (n=83) | P value | Moderate/ severe BPD (n=12) | Nonmoderate/ severe BPD (n=11) | P value | Moderate/ severe BPD (n=9) | Non- moderate/ severe BPD (n=72) | P value |
|---------------------|-----------------------------|----------------------------------|---------|-----------------------------|-----------------------------|---------|-----------------------------|----------------------------------|---------|
| RDW at birth        | 15.8 (15.5–16.7) | 16.2 (15.5–16.7) | .650 | 15.8 (15.2–16.2) | 15.8 (15.5–16.3) | .797 | 15.8 (15.4–19.2) | 16.2 (15.5–16.8) | .964 |
| RDW at D7           | 17.2 (16.1–20.0) | 17.0 (16.1–18.2) | .419 | 17.7 (16.3–21.2) | 16.5 (15.9–18.7) | .174 | 16.9 (15.7–20.3) | 17.1 (16.1–18.3) | .792 |
| RDW at D28          | 17.9 (16.7–18.9) | 17.7 (16.4–18.8) | .634 | 18.1 (17.2–19.0) | 17.6 (16.6–18.5) | .477 | 16.9 (16.7–18.6) | 17.6 (16.3–18.8) | .748 |
| RDW birth-D7        | 0.9 (0.6–2.3) | 0.8 (0.2–1.5) | .289 | 1.5 (0.6–5.3) | 1.0 (0.3–2.6) | .217 | 0.7 (0.2–1.0) | 0.8 (0.1–1.5) | .798 |

BPD = bronchopulmonary dysplasia, RDW = red blood cell distribution width.

Data were presented as median (interquartile range).
preterm infants born at less than 34 weeks’ GA than in neonates of later GA. The 2 previous studies reported that mean RDW was the highest at 32 to 34 weeks’ gestation and suggested that it was secondary to active erythropoiesis in the third trimester. In the present study of preterm infants with a lower GA than in the previous studies, the RDW levels at birth were higher in infants born at 28 to 29 weeks’ GA than in infants born at <28 weeks’ GA. Alur et al reported similar results in that the RDW in the 26 to 31 weeks’ GA group was higher than that in the <26 weeks’ GA group, although the difference was not statistically significant. We suggested that RDW increased gradually, peaked at 32 to 34 weeks’ GA, and decreased again. The RDW values in both the <28 weeks’ and ≥28 weeks’ GA groups increased during the first week of life, while the gap in RDW levels between the groups decreased. Christensen et al explained that the increase in RDW in the early days of life in preterm infants was secondary to previous RBC transfusion. However, the RDW showed similar trends in the present study, although the infants who received RBC transfusions in the first week of life were excluded. We cautiously speculate that preterm infants are vulnerable to postnatal environments such as inflammation and abrupt cessation of iron transplacental transportation, which introduce ineffective erythropoiesis. Christensen et al suggested that high RDW levels is due to reticulocytosis.

In adult studies, a number of studies have reported the role of RDW as an indicator of severity or predictors of outcome for various diseases including sepsis, respiratory disease, cardiovascular disease, and critical illness. Although the mechanism of RDW increase has not been fully determined, it has been suggested that chronic hypoxia, malnutrition, and inflammation cause an increase in RDW values. Hypoxia induces erythropoietin release, which leads to the release of immature reticulocytes into the circulation. Injured RBCs by inflammation aggravate disease progression by decreasing oxygen transfer to organs and tissues.

RDW values were higher in patients with chronic obstructive pulmonary disease than in healthy people and associated with its severity and outcomes, including mortality and readmission rates. The pathophysiology of BPD and chronic obstructive pulmonary disease are similar. Both result from the impairment of alveolarization/vascularization after inflammation and oxidative stress. Recently, Go et al reported that RDW at D28 can predict BPD and its severity. But they found RDW at birth and at D14 were not related to BPD. Although they performed multivariate logistic regression with RBC transfusion, they did not account for transfusion between D14-D28. RBC transfusion is known to affect RDW levels and can limit its role in the prediction of BPD. In addition, infants with BPD need oxygen or positive pressure ventilation and are likely to get a transfusion. As a result, including infants with transfusion may introduce a confounding factor in the analysis, and this may affect the results. Garofoli et al also showed that the RDW at the 4th week of life was higher in the BPD group than in the non-BPD group, whereas there was no difference in RDW within the first 3 days of life between the groups. However, they did not consider GA. In contrast, Doğan found that RDW at birth was related to BPD, which contradicts the findings of the other 2 studies. However, their findings were strongly different from previous studies, with high RDW levels being associated with an odds ratio of 11.9. In the present study, the RDW values at birth, D7, D28, and the change between birth to D7 were not related to the development of BPD. There was no difference even when we evaluated the data further by dividing it into 2 groups of GA, considering the difference in RDW levels and incidence of BPD in accordance to GA.

There were some limitations to this study. First, morbidities such as necrotizing enterocolitis and patent ductus arteriosus were not evaluated due to their complex relationships to BPD over time. However, we investigated whether RDW could be a predictor of BPD independently of other morbidities in a clinical setting. Second, the babies who received RBC transfusion between at birth and D7 were excluded in order to prevent them from affecting RDW values. However, this may have induced a selection bias in which more vulnerable babies were excluded. Further, while we excluded infants who got transfusions in early days of life, we still have this limitation regarding RDW at D28.

The strength of this study was in evaluating changes of RDW in early days of life as well as RDW levels at birth and D7, and their relationships with BPD presentation, even though our study did not show any statistical significance in our results. We also divided our patients into 2 groups born <28 weeks’ GA and ≥28 weeks’ GA, because both RDW levels and incidence of BPD can be quite different according to GA.

In conclusion, RDW values vary in accordance with GA and they change during the newborn period. RDW values at birth were higher in infants born at 28 to 29 weeks’ GA than in those

### Table 4

| Multivariable association between RDW and BPD. | Univariate | | | multivariable | | | |
|---|---|---|---|---|---|---|---|
| | OR | 95% CI | P value | OR | 95% CI | P value |
| BPD | | | | | | | |
| RDW at birth | 1.145 | 0.851–1.541 | .37 | 0.964 | 0.683–1.361 | .381 |
| RDW at D7 | 1.165 | 0.964–1.407 | .113 | 1.054 | 0.835–1.332 | .658 |
| RDW at D28 | 1.153 | 0.905–1.469 | .248 | 1.031 | 0.791–1.344 | .822 |
| RDW birth-D7 | 1.187 | 0.931–1.514 | .167 | 1.119 | 0.829–1.511 | .462 |
| Moderate/severe BPD | | | | | | | |
| RDW at birth | 1.093 | 0.788–1.515 | .595 | 0.868 | 0.565–1.333 | .518 |
| RDW at D7 | 1.188 | 0.989–1.426 | .065 | 0.88 | 0.683–1.134 | .323 |
| RDW at D28 | 1.059 | 0.790–1.420 | .702 | 0.937 | 0.629–1.394 | .747 |
| RDW birth-D7 | 1.261 | 1.002–1.587 | .048 | 0.854 | 0.591–1.235 | .402 |

RDW = bronchopulmonary dysplasia, CI = confidence interval, D28 = twenty-eighth day of life, D7 = seventh day of life, OR = odds ratio, RDW = red cell distribution width.

* Adjusted by birth weight, small for gestational age, and white blood cell count.
born at <28 weeks’ GA. RDW values of both groups increased in the first week of life and remained similar during the first month of life. More care is necessary when assessing RDW values in relation to neonatal morbidity. The RDW values in the first month of life were not associated with BPD development independent of severity in this study. Thus, the usefulness of RDW as a predictor of BPD remains unknown and requires further large-scale studies.

Author contributions

Conceptualization: Seong Hee Oh, Chan-Hoo Park.

Data curation: Seong Hee Oh, Hyun-Jeong Do, Jae Young Cho.

Formal analysis: Seong Hee Oh, Hyun-Jeong Do, Jae Young Cho.

Investigation: Hyun-Jeong Do, Ji Sook Park, Jae Young Cho.

Methodology: Seong Hee Oh, Chan-Hoo Park.

Visualization: Seong Hee Oh, Ji Sook Park.

Writing – original draft: Seong Hee Oh.

Writing – review & editing: Seong Hee Oh, Hyun-Jeong Do, Ji Sook Park, Jae Young Cho, Chan-Hoo Park.

References

[1] Roberts GT, El Badawi SB. Red blood cell distribution width index in some hematologic diseases. Am J Clin Pathol 1985;83:222–6.

[2] Hu ZD, Chen Y, Zhang L, et al. Red blood cell distribution width is a potential index to assess the disease activity of systemic lupus erythematosus. Clin Chim Acta 2013;425:202–5.

[3] Perlstein TS, Weuve J, Pfeffer MA, Beckman JA. Red blood cell distribution width and mortality risk in a community-based prospective cohort. Arch Intern Med 2009;169:588–94.

[4] Ellahomy DM, El-Mekkawy MS, Farag MM. A study of red cell distribution width in neonatal sepsis. Pediatr Emerg Care 2020;36:378–83.

[5] Ytas JW, Horrow JC, Horne BD. Persistent increase in red cell size distribution width after acute diseases: a biomarker of hypoxemia? Clin Chim Acta 2015;448:107–17.

[6] Mohindra R, Mishra U, Mathew R, Negi NS. Red cell distribution width index (RDW) as a predictor of severity of acute ischemic stroke: a correlation study. Adv J Emerg Med 2020;4:824.

[7] Gravito-Soares M, Gravito-Soares E, Gomes D, Almeida N, Tome L. Red cell distribution width and red cell distribution width to total serum calcium ratio as major predictors of severity and mortality in acute pancreatitis. BMC Gastroenterol 2018;18:108.

[8] Epstein D, Nasser R, Mashiachi T, Azzam ZS, Berger G. Increased red cell distribution width: a novel predictor of adverse outcome in patients hospitalized due to acute exacerbation of chronic obstructive pulmonary disease. Respir Med 2018;136:1–7.

[9] Lee JH, Chung HJ, Kim K, et al. Red cell distribution width as a prognostic marker in patients with community-acquired pneumonia. Am J Emerg Med 2013;31:72–9.

[10] Braun E, Domany E, Keng Y, Mazor Y, Makhoul BF, Azzam ZS. Elevated red cell distribution width predicts poor outcome in young patients with community acquired pneumonia. Crit Care 2011;15:R194.

[11] Patel KV, Seema RD, Ferrucci L, et al. Red cell distribution width and mortality in older adults: a meta-analysis. J Gerontol A Biol Sci Med Sci 2010;65:258–65.

[12] Kim S, Lee K, Kim I, Jung S, Kim MJ. Red cell distribution width and early mortality in elderly patients with severe sepsis and septic shock. Clin Exp Emerg Med 2015;2:157–61.

[13] Jo YH, Kim K, Lee JH, et al. Red cell distribution width is a prognostic factor in severe sepsis and septic shock. Am J Emerg Med 2013;31:545–8.

[14] Lorente I, Martin MM, Abreu-Gonzalez P, et al. Red blood cell distribution width during the first week is associated with severity and mortality in septic patients. PLoS One 2014;9:e105436.

[15] Wei S, Cui H, Zhang S, Zhang A, Zhang Y, Jiang S. Red blood cell distribution width predicts postoperative death of infective endocarditis. Int Heart J 2020;61:524–30.

[16] Pinho J, Marques EF, Freitas E, et al. Red cell distribution width as a predictor of 1-year survival in ischemic stroke patients treated with intravenous thrombolysis. Thromb Res 2018;164:4–8.

[17] Al-Najar Y, Goode KM, Zhang J, Cleland JG, Clark AL. Red cell distribution width: an expensive and powerful prognostic marker in heart failure. Eur J Heart Fail 2009;11:1155–62.

[18] Sachdev A, Small A, Khowar A, Gupta N, Gupta D, Chugh P. Outcome prediction value of red cell distribution width in critically-ill children. Indian Pediatr 2018;55:414–6.

[19] Zuk M, Migdal A, Dominczak J, Brzezinska-Rajsz G. Usefulness of red cell distribution width (RDW) in the assessment of children with pulmonary arterial hypertension (PAH). Pediatr Cardiol 2019;40:820–6.

[20] Ramby AL, Goodman DM, Wald EL, Weiss SL. Red blood cell distribution width as a pragmatic marker for outcome in pediatric critical illness. PLoS One 2015;10:e0129258.

[21] Polat V, Işcan S, Eti M, et al. Red cell distribution width as a prognostic indicator in pediatric heart disease and after surgery. Biomed Res Int 2014;2014:681679.

[22] Martin SL, Desai S, Nanavati R, Colah RB, Ghosh K, Mukherjee MB. Red cell distribution width and its association with mortality in neonatal sepsis. J Matern Fetal Neonatal Med 2019;32:1925–30.

[23] Sahgeh S, Sepidarkish M, Mohseni SO, Movahedian A, Mosayebi Z. Red cell distribution width as a predictor of persistent pulmonary hypertension of the newborn. Am J Perinatol 2017;34:1442–6.

[24] Özer Bekmez B, Tayman C, Buyukyturya M, Cetinkaya AK, Akar U, Derme T. A promising, novel index in the diagnosis and follow-up of patent ductus arteriosus: red cell distribution width-to-platelet ratio. J Clin Lab Anal 2018;32:e22616.

[25] Garofoli F, Cardelli L, Mazzuccelli I, et al. The red cell distribution width (RDW): value and role in preterm, IUGR (intrauterine growth restricted), full-term infants. Hematology (Amsterdam; Netherlands) 2014;19:365–9.

[26] Ge H, Ohno H, Nollet KE, et al. Red cell distribution width as a predictor for bronchopulmonary dysplasia in premature infants. Sci Rep 2021;11:7221.

[27] Peln D. The role of red cell distribution width as a predictor of bronchopulmonary dysplasia in preterm infants. Haydarpaşa Numune Med J 2019;60:41–5.

[28] Jöbe AH, Bancalari E. Bronchopulmonary dysplasia. Am J Respir Crit Care Med 2001;163:1723–9.

[29] Christensen RD, Yash HM, Henry E, Bennett ST. Red blood cell distribution width: reference intervals for neonates. J Matern Fetal Neonatal Med 2015;28:883–8.

[30] Tonbul A, Tayyem C, Cortal F, Kara S, Tatif MM. Red cell distribution width (RDW) in the newborn: normative data. J Clin Lab Anal 2011;25:422–5.

[31] Desai SA, Martin SL, Nanavati RN, et al. Red cell distribution width (RDW): normative data in Indian Neonates. J Matern Fetal Neonatal Med J 2019;60:119–21.

[32] Alur P, Devapatla SS, Super DM, et al. Impact of race and gestational age on red blood cell indices in very low birth weight infants. Pediatrics 2000;106(2 Pt 1):306–10.

[33] Teterminz KC, Ongen Alpaydin A, Sevici C, et al. red cell distribution width “predict COPD severity?” Rev Port Pneumol (2006) 2015;21:196–201.

[34] Özgel G, Seyhan EC, Özgel MA, Gunluoglu MZ. Red blood cell distribution width in patients with chronic obstructive pulmonary disease and healthy subjects. Archivos de bronconeumología 2017;53:107–13.

[35] Tonelli M, Sacks F, Arnold M, Moye L, Davis B, Pfeffer M. Relation between red blood cell distribution width and cardiovascular event rate in people with coronary disease. Circulation 2008;117:163–8.

[36] Seyhan EC, Özgel MA, Tutar N, Omer I, Uysal A, Altun S. Red blood cell distribution and survival in patients with chronic obstructive pulmonary disease. Copd 2013;10:416–24.

[37] Pierce CN, Larson DF. Inflammatory cytokine inhibition of erythropoiesis in patients implanted with a mechanical circulatory assist device. Perfusion 2005;20:83–90.

[38] Bourbon JR, Boucherat O, Boczkwoski J, Crestani B, Delacourt C. Bronchopulmonary dysplasia and emphysema: in search of common therapeutic targets. Trends Mol Med 2009;15:169–79.

[39] Spadaro S, Taccone FS, Fogagnolo A, et al. The effects of blood transfusion on red blood cell distribution width in critically ill patients: a pilot study. Transfusion 2018;58:1863–9.

[40] Jiang W, Zou Z, Zhao S, et al. Erythrocyte transfusion limits the role of elevated red cell distribution width on predicting cardiac surgery associated acute kidney injury. Cardiol J 2021;28:255–61.