Red Blood Cell Distribution Width as a Marker of Early Onset Neonatal Sepsis: A Hospital Based Analytical Study

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Abstract
Background: Neonatal sepsis is still a significant cause of neonatal morbidity & mortality and its early diagnosis still remains a challenge. Present study aimed to investigate Red cell Distribution Width (RDW) as a marker of early onset neonatal sepsis (EONS).

Material and Methods: A hospital based cross sectional analytical study was conducted at department of paediatrics, at a tertiary care centre of western India, including 55 early onset neonatal sepsis cases and 55 controls. Newborn delivered by spontaneous vaginal delivery with gestational age of 36 weeks or more presenting within 72 hours of delivery with signs & symptoms suggestive of early onset neonatal sepsis were taken as cases and age and weight-matched healthy new-born were taken as control.

Observations: Mean RDW level was significantly higher (p<0.001) in EONS (21.31 ± 3.08 %) as compared to healthy controls (16.23 ± 1.16%). TLC was also significantly higher (p<0.001) in EONS (18.92 ± 8.01x10^3/dl) as compared to controls (12.19 ± 9.29x10^3/dl). The area under the ROC curve for RDW was 0.988 (0.973 – 1.000), which indicated that RDW was a good predictor of EONS (p<0.001). The Critical cut off point was found to be 18.55 using Youden’s index with a sensitivity of 94.55% and specificity of 96.36% for diagnosis of EONS.

Conclusion: Red cell distribution width can be used as a cheap, easy, rapid and accurate marker for rapid identification of early onset neonatal sepsis.

Keywords: Neonatal sepsis, early onset neonatal sepsis, Red cell distribution width, RDW.

Introduction
Neonatal Sepsis is an invasive infection occurring within the first 28 days of life. Despite advances in perinatal & neonatal care, neonatal sepsis is still a significant cause of mortality & morbidity. According to National Neonatal Perinatal Database (NNPD, 2002-03) incidence of neonatal sepsis in India was 30 per 1000 live births and it contributed to 18.6% of neonatal deaths, next to only perinatal asphyxia (28.8%) and prematurity (26.3%) [1]. Neonatal sepsis is defined as systemic inflammatory response syndrome in the presence of or as a result of suspected or proven infection [2]. It encompasses various systemic infections of
the newborn such as Septicemia, Meningitis, Pneumonia, Arthritis, Osteomyelitis, and Urinary tract infections.

Early onset neonatal sepsis (EONS) presents within the first 72 hours of life. Infants with EONS usually present with respiratory distress and pneumonia. The source of infection is generally the maternal genital tract[3]. EONS constituted 67% of all neonatal sepsis cases in India[1]. The micro-organisms most commonly associated include Group B Streptococcus, E. coli, Haemophilus influenzae and Listeria monocytogenes[4].

Risk factors associated with an increased risk of EONS include Low birth weight or prematurity, febrile illness in the mother with evidence of bacterial infection within 2 weeks prior to delivery, foul smelling liquor, rupture of membranes >24 hours, single unclean or >3 sterile vaginal examination(s) during labour, prolonged labour and perinatal asphyxia (Apgar score < 4 at 1 minute). Immature immune defences in newborns, including low circulating levels of immunoglobulins, decreased number of T lymphocytes and neutrophils, and functionally impaired cytotoxic activity in leukocytes[5-7] are involved in pathogenesis of neonatal sepsis.

Diagnosis of early onset neonatal sepsis remains a challenge for neonatal health care providers. The gold standard for detecting bacterial sepsis is blood culture, but its positivity ranges from as low as 8% to 73%[8-10] in the diagnosis of potential neonatal sepsis and it has limitation of 24 to 48 hour assay time. Low sensitivity of culture for identifying neonatal sepsis[5,11] and limited accuracy and reliability of haematological markers have led to development of scoring system for diagnosis of EONS. These scoring systems include clinical and laboratory findings (total WBC counts, absolute number of neutrophils, immature/total neutrophil ratio) and acute phase reactants (CRP and procalcitonin) in different combinations[5-9].

Red blood cell distribution width (RDW) is a parameter reflecting the heterogeneity of the peripheral red blood volume and is usually expressed with RDW-coefficient of variation (RDW-CV). Limited information is available about the association between RDW and EONS[12]. This study, aimed to compare RDW levels in healthy new-born and those with EONS in order to investigate the potential role of RDW as a marker of EONS.

**Material and Methods**

This study was a hospital based cross sectional analytical study conducted at department of paediatric medicine, at a tertiary care centre of western India,. A total of 55 early onset neonatal sepsis cases and 55 controls presenting to the hospital over the period of sep-2017 to nov-2018 were taken into the study. Newborn delivered by spontaneous vaginal delivery with gestational age of 36 weeks or more presenting to the hospital within 72 hours of delivery with signs & symptoms suggestive of early onset clinical infection or pneumonia with presence of foul smelling liquor or least two of the following risk factors - Febrile illness in the mother with evidence of bacterial infection within 2 weeks prior to delivery, rupture of membranes >24 hours, single unclean or >3 sterile vaginal examination(s) during labour.

Prolonged labour (sum of 1st and 2nd stage of labour > 24 hrs), Perinatal asphyxia (apgar score <4 at 1 minute). Age and body weight-matched healthy new-born in postnatal ward or those brought to the outpatient clinics for neonatal screening program within 72 hours of delivery were taken as control. Very-low-birth (<2500 g) infants, those with perinatal asphyxia, meconium aspiration syndrome, congenital malformations, congenital infections associated with the TORCH complex, metabolic disease, Rh or ABO isoimmunisation or those delivered by Cesarean section were excluded from the study.

Eligible selected subjects were subjected to a detailed history, clinical examination & laboratory evaluation. Peripheral blood samples were drawn...
during their first hospital visits and CBC, and blood typing was determined.
Diagnosis of EONS was made according to following criteria:
1) Clinical signs consistent with infection based on the Töllner score or
2) Culture-positivity detected in blood, urine and cerebrospinal fluid samples or
3) Signs of pneumonia on chest X-ray.
Clinical parameters used in the Töllner scoring system include discoloration of skin, isordered Peripheral Circulation, Hypotonia, Bradycardia, Apnea and Respiratory Distress, Hepatomegaly, Abdominal Distension, Abnormal Laboratory Parameters (Left-Shift In WBC Counts, Thrombocytopenia) and Metabolic Acidosis. For each parameter one point is assigned. Patients were categorized clinical sepsis (≥10 pts), suspicion of sepsis (5-10 pts) and normal (≤5 pts) new-born were categorized\[7\].

**Statistical Analysis:** Nominal / categorical variables were summarized as frequency and percentage and analyzed using Chi square test. Continuous variables were summarized as mean and standard deviation and were analyzed using independent sample t test. Multivariate logistic regression analysis was done to determine the independent predictors of EONS. ROC curve was used to determine the role of RDW for diagnosis of EONS and to determine the critical cutoff using Youden index. A p value < 0.05 was taken as statistically significant. All statistical analysis was done using SPSS trial version 20 statistical software.

**Observation**
The baseline characteristics of neonatal sepsis cases and healthy controls were comparable (Table 1). Mean RDW level was found to be significantly higher (p<0.001) in EONS cases (21.31 ± 3.08 %) as compared to healthy controls (16.23 ± 1.16%). TLC was also found to be significantly higher (p<0.001) in EONS (18.92 ± 8.01x10^3/dl) as compared to healthy controls (12.19 ± 9.29x10^3/dl). (Table 2). Multivariate logistic regression analysis showed that RDW (OR = 50.39) and TLC (OR= 1.204) were independent predictors of EONS (Table 3). The area under the ROC curve for RDW was 0.988 (0.973 – 1.000), which indicated that RDW was a good predictor of EONS (p<0.001). The Critical cut off point was found to be 18.55 using Youden’s index which indicates that RDW ≥18.55 was diagnostic of EONS (Figure 1). At cutoff of 18.5, RDW had 94.55% sensitivity and 96.36% specificity for diagnosis of EONS (Table 4).

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### Table 1: Baseline characteristics of study groups

|                                | Case (N=55)   | Control (N=55) | p value |
|--------------------------------|---------------|----------------|---------|
| **Age in years: (mean ± SD)**  | 26.56 ± 19.0  | 26.07 ± 12.31  | 0.873   |
| **Gender – Male**              | 37 (%)        | 34 (%)         | 0.690   |
| **Female**                     | 18 (%)        | 21 (%)         |         |
| **Religion – Hindu**           | 47 (%)        | 49 (%)         | 0.775   |
| **Muslim**                     | 8 (%)         | 6 (%)          |         |
| **Gestational age in weeks**   | 38.20 ± 1.16  | 37.89 ± 0.90   | 0.120   |
| **Birth weight in Kg**         | 2.74 ± 0.18   | 2.67 ± 0.39    | 0.267   |
| **Hemoglobin in gm%**          | 16.26 ± 2.39  | 15.56 ± 1.87   | 0.089   |

### Table 2: Comparison of Blood parameters among study groups

|                  | Case           | Control        | p value   |
|------------------|----------------|----------------|-----------|
| **TLC (x10^3/dl)**| 18.92 ± 8.01   | 12.19 ± 9.29   | <0.001 (S) |
| **Platelet count (x10^5/dl)**| 2.56 ± 1.16     | 2.99 ± 1.24    | 0.062     |
| **RDW (%)**      | 21.31 ± 3.08   | 16.23 ± 1.16   | P<0.001 (S) |

TLC- Total leucocyte count; RDW- Red cell distribution width
Table 3: Multivariate logistic regression analysis to determine predictors of EONS

| Variables          | Odds ratio | 95% C.I. | Coefficient (B) | SE  | P value |
|--------------------|------------|----------|-----------------|-----|---------|
| Age                | 0.942      | .858     | 1.034           | -0.060 | 0.048  | 0.209 |
| Sex(Male)          | .069       | .001     | 3.237           | -2.677 | 1.965  | 0.173 |
| Religion(Muslim)   | .007       | .000     | 3.254           | -4.973 | 3.139  | 0.113 |
| Gestational Age    | 1.793      | .253     | 12.689          | .584  | .998   | 0.559 |
| Birth Weight       | 8.14       | .006     | 46.6            | 4.473 | 4.407  | 0.311 |
| Hemoglobin         | 2.812      | .928     | 7.689           | 0.934 | .513   | 0.144 |
| TLC                | 1.204      | 1.016    | 1.427           | .185  | .087   | 0.033*|
| Platelet count     | .382       | .091     | 1.598           | -.963 | .730   | 0.187 |
| RDW                | 50.39      | 3.239    | 784.113         | 3.920 | 1.400  | 0.005*|

Figure 1: ROC curve of RDW for diagnosis of EONS

Table 4: Diagnostic parameters of RDW for diagnosis of EONS

| Parameter     | Value     | 95% CI          |
|---------------|-----------|-----------------|
| Sensitivity   | 94.55%    | 85.15 – 98.13%  |
| Specificity   | 96.36%    | 87.46 – 98.98%  |
| PPV           | 96.3%     | 87.46 – 98.98%  |
| NPV           | 94.64%    | 85.39 – 98.16%  |
| Diagnostic accuracy | 95.45%   | 89.8 – 98.04%   |

Discussion

Neonatal sepsis with its high mortality rate, still remains a diagnostic and treatment challenge for the neonatal health care providers. An early diagnosis of neonatal septicemia helps in instituting antibiotic therapy at the earliest time, thereby reducing the mortality rates and also avoids the unnecessary treatment of a non infected neonate. Mortality rate are high in the developing countries because of the lack of early diagnosis and identification of high risk cases\(^\text{[13]}\). Neonatal sepsis also increases the length of hospital stay and cost of treatment especially in developing countries with insufficient neonatal intensive care facilities and personnel. Blood culture which is gold standard for diagnosis of neonatal sepsis not
only takes time, but it is also complicated, with a 
low yield[14]. For this reason, a number of 
biochemical markers have been tested for accurate 
diagnosis of sepsis in the shortest time[15]. 
In this study, red cell distribution width was tested 
as an early inexpensive and easily available 
biomarker for diagnosis of neonatal sepsis. Mean 
age of cases was 26.56±19.00 hours and that of 
controls was 26.07±12.31 hours. Cosar H et al[15] 
reported similar findings with mean age of cases 
being 1.98±0.9 days and that of controls being 
1.87±0.92 Days. In the present study 67.27% of 
cases were male, similar male preponderance was 
found by cosarh et al[15] and Betty chacko et al[16]. 
Medhat A Saleh et al[17] however observed 
number of females to be more than male. 
In present study the mean RDW level was 
significantly higher in neonatal sepsis cases 
(21.31±3.08%) as compared controls 
(16.23±1.16%). Jianping et al[18], also found mean 
RDW level was significantly higher than sepsis 
group. This can be explained by the fact that 
inflammation causes an increase of neurohormone 
and endocrine hormone in the body including 
noradrenaline, angiotensin I and other angiotensin 
levels and renal ischemia. These neurotransmitters 
can stimulate red blood cell proliferation through 
increased secretion of erythropoietin (EPO) 
causing increase in RDW and inflammatory 
factors may affect marrow hematopoietic function 
and iron metabolism in the body to cause increase 
in RDW value[19].

In this study, increased RDW was found in in 
term and near term infants with EONS. RDW is a 
low-priced arithmetical index and is part of a 
standard complete blood count. RDW is quickly 
obtained, and also does not require additional 
costs and can be provided easily. Comparative 
studies investigating the correlations between 
sepsis, septic shock and RDW have been more 
frequently performed with adult population. Chan 
Ho Kim et al[20], in a study of 329 patients, found 
higher mortality rates in patients with higher 
RDW. In study by Ku, Nam Su; Kim et al[21] 
including 566 adult patients, RDW was indicated 
as a potentially independent prognostic factor for 
28 day-mortality in patients with sepsis and septic shock[21]. 
Christensen RD, Yaish HM et al[22] investigated 
reference ranges of RDW for newborn, and 
detected lower and upper limits of normal at birth, 
for term and late-preterm infants as 15.5% and 
20%, respectively. However, in premature infants, 
upper limit of RDW was higher (23%)[23]. In 
present study, RDW (21.31±3.08%) measured 
within 4 days of the postnatal period was similar 
to the lower limit of normal reference range of 
term and late-term infants in the above-mentioned 
study.

Multivariate logistic regression analysis revealed 
RDW (p=0.005) and TLC (P=0.033) were 
independent predictors of EONS. Higher RDW 
strongly suggested of EONS (OR =50.39) and 
TLC (OR=1.204) was a weak indicator. Area 
under the ROC curve for RDW was found to be 
0.988 (0.973 – 1.000) indicating that RDW was a 
good predictor of EONS. The Critical cut off point 
was found to be 18.55 using Youden index which 
indicates that RDW ≥18.55 is diagnostic of 
EONS.

The sensitivity and specificity of RDW to detect 
EONS in cases included on the basis of clinical 
signs and culture positivity was 94.50% and 
96.36% respectively. Abdullah ST et al[19] in a 
study on 112 newborns, found that at cut off of 
RDW 14.3% had sensitivity and specificity of 
85% and 100% respectively for diagnosis of 
sepsis.

Though the mechanism of increased RDW in 
Sepsis is not known, higher RDW levels 
demonstrate its association with inflammatory 
processes. It has been detected that markers of 
inflammation including RDW-associated 
interleukin-6 (IL-6), tumour necrosis factor-alfa 
(TNF-α) and proinflammatory cytokines suppress 
maturity process of RBC and increase their 
half-lives with resultant rise in RDW levels[10,23]
Conclusion
RDW was observably higher in new-born with EONS. It is a hematological index estimated within the standard CBC profile and can be easily and rapidly estimated without additional cost. Further studies with preferably prospective design and wider clinical profile could help establish role of RDW alone or in combination other haematological score for early and rapid diagnosis of EONS.

Conflict of interest: None

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