Cerebral Hemodynamics in Stable Preterm Infants Before and After Packed Cell Transfusion

Chinmay Chetan
Bharati Vidyapeeth Deemed University

Nyein Zaw
Women and Children Hospital, Taunggyi

Pradeep Suryawanshi (drpradeepsuryawanshi@gmail.com)
Bharati Vidyapeeth Deemed University

Nishant Banait
Bharati Vidyapeeth Deemed University

Prince Pareek
Bharati Vidyapeeth Deemed University

Sujata Deshpande
Bharati Vidyapeeth Deemed University

Bhvyaa Gupta
Bharati Vidyapeeth Deemed University

Reema Garegrat
Bharati Vidyapeeth Deemed University

Arjun Verma
Bharati Vidyapeeth Deemed University

Research Article

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Abstract

**BACKGROUND** In a year around 3.5 million preterm deliveries occur in India alone. Some of these babies will require packed cell volume (PCV) transfusion. There is a paucity of robust data on effect of blood transfusions on the cerebral hemodynamic from India. This study was done to see the effect of PCV transfusion on blood flow velocities and resistive index (RI) of anterior cerebral artery (ACA) in stable preterm infants.

**METHOD** A prospective observational study was conducted in a tertiary care hospital in Pune, India. All stable preterm infants (<37 weeks) receiving PCV transfusion were enrolled. USG Doppler study of ACA was done before and after PCV transfusion. Peak systolic velocity (PSV), end-diastolic velocity (EDV) and RI were measured pre and post PCV transfusion.

**RESULTS** Thirty infants were included in the study, with median gestation age of 28.8 [interquartile range \{IQR\}, 27-30.55] weeks and median birth weight of 970 [interquartile range \{IQR\}, 869.5-1190] grams. There was a significant decrease in PSV pre and post PCV transfusion - 58.46 (±18.44) cm/sec and 46.34 (±13.93) cm/sec respectively (p value <0.001). Changes in RI and EDV were non-significant.

**CONCLUSION** PCV transfusion significantly decreased PSV, reflecting improved cerebral oxygenation, and decreased cardiac output after correction of anaemia. Laboratory threshold for PCV transfusion in stable preterm infants are not known. USG Doppler study has the potential to provide one of the objective criteria for PCV transfusion in these infants though large scale randomized controlled trials are needed to prove its efficacy.

Background

An estimated 15 million babies are born preterm every year, globally [1]. The largest number of preterm births occurs in India [2]. Anaemia is a frequent complication of prematurity and is known to compromise the neurological development of the preterm infants. Many cause lead to anaemia in these babies. Firstly, premature infants do not complete the third trimester of gestation and it is at this stage the maximum iron transport occurs through the placenta. Secondly, at this stage of development, erythropoiesis mainly occurs in the liver and the bone marrow. The liver is less responsive to anemia in stimulating an erythropoietin response. In addition, preterm infants undergo frequent blood tests in neonatal intensive care units (NICU). All these mechanisms of, rapid growth, multiple blood samplings and insufficient erythropoiesis contribute to anemia and its severity. Blood transfusions are common in neonatal units with the intention to maintain optimal tissue oxygenation and promote growth. [3, 4].

One of the principal objectives of blood transfusion in preterm infants is to prevent impaired tissue oxygenation of vital organs and brain.[5]. The benefits of blood transfusion in stable preterm infants with respect to neurological outcomes are not clear.
The main variables affecting cerebral oxygen delivery are cerebral blood flow and haemoglobin concentration [6]. Cerebral blood flow can be measured at the bedside by using non-invasive Doppler ultrasound scan of contributing arteries. The internal carotid artery, basilar artery, anterior cerebral artery and lenticulostriate arteries can be easily visualized with color Doppler imaging [7]. Blood flow can be evaluated by measuring peak systolic velocity (PSV), end-diastolic velocity (EDV) and calculating resistive index (RI). RI measures vascular resistance and is defined as (PSV – EDV) / PSV [8]. A high RI corresponds to vasoconstriction and low blood flow velocity, whereas a low RI is related to vasodilation and high blood flow velocity [9].

Determining peak velocity of systolic blood flow in the middle cerebral artery as an indicator of the severity of anemia is well-documented in fetuses [10]. Researchers have shown that blood transfusions improve cerebral oxygen supply and induce a decrease in cerebral blood flow velocity, by using near infrared spectroscopy (NIRS) and cerebral doppler ultrasonography [11–13]. But there is paucity of robust data on effect of blood transfusions on the cerebral hemodynamic from India. Our study is aimed to document the changes in cerebral blood flow after packed cell transfusion in stable preterm infants.

**Methods**

A prospective observational study was done from April 2018 to December 2019, in a tertiary level NICU at Pune, India. All stable preterm infants receiving packed cell transfusions for clinical indications and a haemoglobin value of less than 10 gm/dl during the study period were enrolled after obtaining written informed consent from parents. Prior ethics approval was taken from the Bharati Vidyapeeth Medical College Institutional ethics committee before starting the study. Preterm infants needing cardio-respiratory support, with major congenital malformations and genetic syndromes were excluded. Transcranial colour Doppler ultrasonography was done 1 hour before packed cell transfusion and 24 hours after post transfusion by neonatologist who was trained in cranial ultrasonography using a SIEMENS machine (Acuson X 300, SIEMENS Medical Solution) with neonatal probe (5–10 Hz transducer). Doppler imaging of the anterior cerebral artery (ACA) was through the anterior fontanelle in the sagittal plane (Fig. 1). Pulse doppler was done to measure PSV and EDV (Fig. 2). RI was then calculated using the formula: RI= (PSV-EDV) / PSV. All measurements were done in thermo-neutral environment ensuring normal body temperature without any pressure provocation in quite infants, using oral sucrose as pacifier with continuous monitoring of oxygen saturation and vitals. Three measurements were recorded each time and mean was calculated. Protocols followed were according to institutional guidelines. Statistical analysis was done using SPSS software version 25.0. Paired t test was used to test the mean difference between RI, PSV and EDV. Throughout the results 5% level of significance was used. All results are shown with 95% confidence interval, with p-value of less than 0.05 been considered significant.

**Results**
Thirty infants were included in the study, with median gestation age of 28.8 [interquartile range {IQR}, 27-30.55] weeks, median birth weight of 970 [interquartile range {IQR}, 869.5–1190] grams. Pre-transfusion median haemoglobin was 8.1 [interquartile range {IQR}, 7.1–8.7] g/dl (Table 1). Pre and post transfusion mean RI were 0.83 (± 0.07) and 0.82 (± 0.07) respectively (p value 0.17). Pre and post transfusion PSV were 58.46 (± 18.44) cm/sec and 46.34 (± 13.93) cm/sec respectively (p value < 0.001) and pre and post transfusion EDV were 9.83 (± 6.64) cm/sec and 8.47 (± 5.15) cm/sec (p value 0.21) (Table 2).

Table 1

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<th>Baseline characteristics of the population</th>
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<td>Birth weight (Grams)</td>
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<td>Pre transfusion Hb (g/dl)</td>
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Table 2

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<tr>
<th>Pre and post PCV transfusion RI, PSV, EDV</th>
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<tr>
<td>Pre PCV</td>
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<td>RI</td>
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<td>0.83 (0.07)</td>
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<tr>
<td>PSV (cm/sec)</td>
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<td>EDV (cm/sec)</td>
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Discussion

In this study we found a significant decrease in PSV post transfusion but changes in RI and EDV were non-significant. These results are in line with the previously published studies, which showed that blood transfusion decreases the cerebral blood flow velocity [11–13].

Anaemia may lead to increase in cardiac output and cerebral vasodilation, as a compensatory mechanism to increase the oxygenation of brain parenchymal tissue. Previous studies have documented that blood transfusion decrease cardiac output and heart rate. [14]. The decrease cerebral blood flow velocity effect of transfusion can be explained by the reduced cardiac output after the correction of anaemia. Decrease in systolic velocity, may also reflect cerebral vasoconstriction due to improved
cerebral oxygenation. Cerebral blood flow is also affected by fetal hemoglobin concentration [15]. On transfusing adult hemoglobin, oxygen delivery capacity of blood to the tissues increases, further resulting in decrease of cerebral blood flow.

Caution should be excised as flow should not be equated with blood velocity measured via Doppler study. Cerebral blood flow velocity also depends on the cross-section area of the vessel. Unfortunately, diameter of the cerebral blood vessels cannot be measured accurately by the currently available USG machines, therefore we have to rely on the cerebral blood flow velocity to estimate the cerebral blood flow.

Change in only PSV, with no significant change in RI and EDV, may represent ongoing adaptive mechanism after PCV transfusion. Also, PSV and EDV depends on the angle of insonation, whereas RI being a ratio, is not affected by the angle of insonation. [16]. The difference in PSV, could be due to the intra-observer bias or due to change of angle of insonation, between the pre and post PCV transfusion velocity readings. However, we tried to minimize the bias, by standardizing the area, where Doppler was acquired. Angle of insonation was always less than 15 degrees. All readings were measured by same individual.

In spite of many trials and clinical guidelines the criteria for PCV transfusion in preterm babies are not very clear. Frequently, preterm babies receive packed cell transfusion based on clinical signs of inadequate weight gain, tachycardia, tachypnoea, or persistent oxygen requirement. [17] Liberal transfusion increases the risk of transmission of infection, bronchopulmonary dysplasia, necrotising enterocolitis and retinopathy and simultaneously increases the cost of NICU stay [18–20]. On the other hand, restrictive regime may increase the chances of chronic hypoxemic changes, intraparenchymal brain haemorrhage or periventricular leukomalacia [21]. Some studies showed no difference in neonatal mortality or major morbidity when lower haemoglobin threshold was kept for transfusion, but some showed weak evidence of improved long-term outcome with higher threshold. [22, 23, 24].

Researchers have explored the possibility of using cerebral blood flow velocities to define a threshold for transfusion in preterm babies [14]. This approach seems a distant possibility for lack of data from randomised controlled trials.

**Conclusion**

The hemoglobin threshold for the PCV transfusion in stable preterm infants is still elusive, in spite of many trials. This probably reflects the need of defining blood transfusion threshold by considering not only the clinical and laboratory parameters but also haemodynamic parameters. Doppler studies may provide one such vital parameter.

**List Of Abbreviations**

PCV - packed cell volume
RI - resistive index
ACA - anterior cerebral artery
PSV - Peak systolic velocity
EDV - end-diastolic velocity
IQR - interquartile range
NICU - neonatal intensive care units
NIRS - near infrared spectroscopy

Declarations

Ethics approval and consent to participate

Prior ethics approval was taken from the Bharati Vidyapeeth Medical College Institutional ethics committee before starting the study. The study was carried in accordance with the guidelines of Bharati Vidyapeeth Medical College. Informed consent was obtained from parents prior to enrolling infants in the study.

Consent for publication

Not applicable

Availability of data and materials

The datasets used and/or analysed during the current study are available from the corresponding author on reasonable request

Competing interests

The authors declare that they have no competing interests

Funding

Not applicable

Authors' contributions

CC, PS, NNZ: conceived the study; CC, NNZ, BG, RG, AV, PP: did the data collection, analysis and interpretation; CC, NB, SD, PS, NNZ, RG, AV, PP, BG: wrote the first draft which was the read, revised and approved by all the authors. All authors approved the final version of manuscript, and are accountable for all aspects related to the study.
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References


Figures
Figure 1

Identification of ACA (Doppler – Angle of insonation <15°)
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Identification of ACA (Doppler – Angle of insonation <15°)
Figure 2

Calculation of PSV, EDV and RI using Doppler in ACA
Figure 2

Calculation of PSV, EDV and RI using Doppler in ACA