Syndrome of Hyperviscosity in the Neonate
(Literature Review).

by

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Introduction
Sixteen years ago, Chaptal et al. (cited by Mackintosh and Walker, 1973) described 5 infants with dyspnea and convulsions attributable to polycythemia. One year later, in 1959, Wood reported 2 polycythemic infants with dyspnea and convulsions attributable to polycythemia as well and thenceforth attention was called to this disorder and numerous papers have subsequently been published on this subject.

Cases in the literature which dealt with polycythemia appeared as maternal-fetal transfusions (Michael and Mauer, 1961), intrauterine parabiotic syndrome (Naeye, 1963), neonatal respiratory distress associated with a high hematocrit (Danks and Stevens, 1964), cardiopulmonary effects of placental transfusion (Buckels and Usher, 1965) the intrauterine twin-to-twin transfusion syndrome (Singh and Sethi, 1969) and many others. The first report on hyperviscosity in the neonate documented in the Indonesian literature was published by Sukadi and Ali-sjahbana (1974) while ours is still in manuscript.

Gross et al. (1973) defined polycythemia as a venous hematocrit of 65% or greater and it is now recognized that polycythemia may produce symptoms in the newborn. It was Baum in 1966 who first reported the existing relation between hematocrit and blood viscosity in the neonate.

Syndrome of hyperviscosity in the neonate deserves special attention of pediatricians in general and neonatologists in particular as newborn infants are particularly susceptible to developing hyperviscosity and treatment directed to lowering viscosity is promising. Clinical manifestations associated with hyperviscosity

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may not only result in permanent sequelae (Gross et al., 1973) but life-threatening as well.

This paper tries to present a brief review on the many-sided problem of the syndrome of hyperviscosity in the neonate.

**Etiology**

Syndrome of hyperviscosity may arise from increased number of cells (polycythemia or leukemia), increased resistance of cells to deformation (sickle cell disease or hereditary spherocytosis), and elevated plasma viscosity. Increased concentration of fibrinogen or the macroglobulins, IgM or SF19, in the myelomatous process or Waldenstrom’s macroglobulinemia may increase plasma viscosity. Polycythemic hyperviscosity is the most common of all hyperviscosity conditions (Wells, 1970).

Neonatal polycythemia may be due to several factors.

1. Relative intrauterine anoxia may increase erythrocyte production (Walker and Turnbull, 1953) by increasing erythropoietin production (Chaptal et al., 1958 as cited by Gatty et al., 1966; Gross et al., 1973; Gersony, 1973). However, Mackintosh and Walker (1973) suggested that maternal hypoxia and fetal asphyxia at birth increase the blood catecholamines in their respective circulations, assumed to cause more vasoconstriction of the umbilical artery than of the vein, encouraging transfer of blood from placenta to fetus. Mackintosh and Walker (1973) cited Philip et al., (1969) as stating that the hypoxic induction of uterine contractions encourages transfer of blood from placenta to fetus.

2. Small for gestational age and postmaturity. Humbert et al. (1969) reporting on polycythemia in small for gestational age infants attributed to possible chronic hypoxia in utero. Six of the 18 Gross’ (1973) polycythemic infants were small for gestational age. Most of Baum’s (1967) sixteen hyperviscous newborns showed evidences of dysmaturity (cited by Schaffer and Avery, 1971). Fetal hypoxia is often associated with placental insufficiency.


In newborn infants with Down’s syndrome there is evidence that a myeloproliferative disorder of the bone marrow contributes to polycythemia.

transfusion is based on (Gatti et al., 1966):

- High hematocrit and hemoglobin values.
- A low proportion of fetal hemoglobin.
- The demonstration of maternal erythrocytes in the infant's blood by differential agglutination which is most convincing.
- The presence of Beta₂A-globulin (IgA) in the infant's serum.

In twin-to-twin transfusion there is severe anemia in one twin and intense plethora in the other. This occurs only in monochorionic (uniovular) twins where vascular anastomosis between the two parts of the placenta has been found to be invariably present. The vascular shunts may be readily identified with milk or infant formula as the contrast media (Coen and Sutherland, 1970).

5. Delayed clamping of the cord.
Infants whose cords are clamped late have larger blood volumes, higher hematocrits and altered hemodynamics when compared to infants whose cords are clamped early. Usher in 1963 demonstrated that about 40 ml blood is transferred within 15 seconds and an average of 166 ml within 5 minutes after delivery, when the infant is held below the introitus and the cord left undisturbed. Buckels an Usher (1965) found that the capillary hematocrit at 4 hours of age averages 56.4% after early (about 5 seconds after birth), and 75.1% after delayed (5 minutes after birth) cord clamping. These findings suggest that placental transfusion may play an important role during the first minutes of life.

**Physiology and Pathophysiology**

A modified classification of polycythemia by Murray (1966) was put forward by Kontras (1972) as follows:

**I. Relative polycythemia, hemococoncentration.**

**II. Primary polycythemia.**
A. Polycythemia vera.
B. Benign familial polycythemia or erythrocytosis.

**III. Secondary polycythemia.**
A. Insufficient oxygen delivery.
1. Low environmental O₂ (high altitude)
2. Impaired ventilation-pulmonary disease, obesity
3. Cyanotic congenital heart disease
4. Abnormal hemoglobins
   a. Methemoglobinemia (congenital, acquired)
   b. Hemoglobin variants
B. Increases in erythrocyte-stimulating substances (Erythropoietin).
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1. Malignant tissue-renal, cerebellar lesions, hepatic, adrenal.
2. Benign lesions
3. Exogenous stimulation-chemical and physical agents

IV. Neonatal polycythemia.

In this respect we will only deal with the neonatal polycythemia.

Walker and Turnbull (1953) had shown that a fall in the oxygen supply to the fetus during intrauterine life does exert a profound influence on the production of erythrocytes and formation of hemoglobin. Normally, the hemoglobin level rises steadily from 9 gm% at the 10th week to 14 - 15 by the 22nd - 24th week. From the 36th week there is a general tendency for the level to rise so that by the 40th week, the range is from 15 to 18.8 gm and by the 43rd week the mean value is 18.8 gm, the range being 16.8 gm to 20.5 gm. In case of prolonged pregnancy the rise is maintained (Fig. 1).

The erythrocyte count shows a similar pattern. The count rises steadily from 1,420,000 at 10 weeks to 3,280,000 at 25 weeks. From then till 36 weeks the increase is slow. From the 36th week the spread out is in the same way as does the hemoglobin so that by the 40th week the range is from 3,740,000 to 4,940,000 with a mean of 4,350,000. After the 40th week the increase continues (Fig. 2).

During the first few hours after birth, an increase in erythrocyte count, hematocrit and hemoglobin takes place. Usher et al., (1963) explained it as partially resulting from a placental transfusion at the time of delivery with subsequent fluid shift decreasing the plasma volume (cited by Oski and Naiman, 1966).

Gross et al. (1973) proposed the following scheme for the etiology and pathogenesis of symptomatic hyperviscosity states in the neonate. (diagram 1).

Neonatal hyperviscosity is considered to be primarily influenced by erythrocyte number. Increased viscosity causes sludging of flow resulting in impaired tissue oxygenation and consequent acidosis leading to decreased erythrocyte deformability. Tissue hypoxia of the brain, kidneys and lungs leads to clinical signs and symptoms. The high viscosity can alter the cerebrospinal fluid secretion, circulation and absorption, and result in a further rise of intracranial pressure accounting for the eye changes frequently seen on funduscopical examination (Christpın and Darke, 1962). Hypoglycemia is a frequently associated finding in polycythemia but the causal relationship has not been clearly demonstrated yet. Hyperbiliurbinemia may be related to premature destruction of erythrocytes in an abnormal circulation as evidenced by burred and fragmented cells often found in polycythemic
FIG. 1: Hemoglobin levels in cord blood of human fetus in normal pregnancy (Adapted from Walker & Turnbull, 1953).

FIG. 2: Red cell count in cord blood of human fetus in normal pregnancy (Adapted from Walker & Turnbull, 1953).
infants and carries the risk of kernicterus. Thrombocytopenia may be due to consumption of thrombocytes during sludging. Disseminated intravascular coagulation may be another complication of hyperviscosity (Kontras et al., 1970). Transfusion may result in circulatory overload which could be held responsible for the cardiomegaly frequently seen on the chest film, respiratory distress and plethora. The newborn erythrocytes are less deformable contributing to whole blood hyperviscosity.

Normovolemic polycythemia is found in infants who are dysmature or small for gestational age (Humbert et al., 1969). Murray et al., (1963), Replogle and Merrill (1970) and Fouron and Hebert (1973) had demonstrated that in normovolemic polycythemia there is a fall in the systemic cardiac output with increased hematocrit related to the higher viscosity and secondarily to the diminished venous return to the right heart. The pulmonary vascular resistance progressively increases with increasing hematocrit and at a hematocrit of about 70%, a right-to-left shunt appears through the ductus arteriosus and foramen ovale creating a peripheral hypoxia which in turn increases the shunting by causing peripheral vasoconstriction (Fouron and Hebert, 1973; Gersony, 1973). It should probably be regarded as the limit above which clinical symptoms could be anticipated (Mackintosh and Walker, 1973). This value of hematocrit is furthermore a justification for therapeutic phlebotomy in hypervolemic polycythemia (Fouron and Hebert, 1973).

Gregg and Wiggers in 1933 (cited by Murray et al., 1963) had shown an increase in cardiac output in hypervolemic polycythemia (as in transfusion syndromes). Murray et al., (1963) showed that the cardiac output is significantly higher in hypervolemia than in normovolemia in all hematocrit ranges compared (Fig. 3).

Cardiac output has a significant inverse correlation to hematocrit in both normovolemic and hypervolemic polycythemia. Cardiac output is approximately twice as high in hypervolemic as in normovolemic polycythemia at any comparable hematocrit.

However, whether or not the cardiac output is increased in hypervolemic polycythemia depends upon the hematocrit under consideration; for example, inspection of fig. 3 shows that cardiac output at a hypervolemic hematocrit of 60% is higher while cardiac output at a hypervolemic hematocrit of 70% is lower than the cardiac output at a normovolemic hematocrit of 45%. A decreased cardiac output with a consequent reduction in tissue blood flow leads to peripheral hypoxia. Replogle and Merrill (1970) stated that the incre-
DIAGRAM 1: Pathogenesis of symptomatic hyperviscosity states in neonate.

INTRAUTERINE ANOXIA (ERYTHROPOIETIN) → TRANSFUSION (FLUID LOSS) → POLYCYTHEMIA → HYPERDYNAMIC STATE → CARDIOMEGALY

RESPIRATORY DISTRESS → PLETHORA

PLASMA PROTEINS → HYPERVISCOSITY → SLUDGING → OLIGURIA

CYANOSIS → HYPOGLYCEMIA → BILIRUBIN INCREASED

PLATELETS DECREASED

NEWBORN RED BLOOD CELL → DEFORMABILITY

ACIDOSIS

HYPOXIA

FIG. 3: Cardiac output, hematocrit curve (Adapted from Murray et al., 1963)

CARDIAC OUTPUT
ML./KG./MIN.

500
400
300
200
100

10 20 30 40 50 60 70 80

HEMATOCRIT, PERCENT
ased oxygen carrying capacity of the erythrocyte-rich blood does not compensate for the reduced cardiac output associated with polycythemia.

Viscosity-Hematocrit and Viscosity-Shear Rate Relations

There is virtually a straight line relation between viscosity and hematocrit over the normal range, but at hematocrit values greater than 65% there is a progressively larger increase in viscosity for each unit change in hematocrit (Baum, 1966; Mackintosh and Walker, 1973) as illustrated in Fig. 4.

At high hematocrit levels minor increases in hematocrit produce marked increases in viscosity.

The viscosity of blood is not fixed but varies with the rate of shear (velocity gradient) and the dimension of the conducting vessel. At low rates of shear, there is a relatively higher viscosity value for a given hematocrit and the curve approximates a linear function. At high shear rates, the curve moves down and to the right with lower values for viscosity at hematocrit levels comparable to those plotted at low rates of shear (Wells and Merrill, 1961) as shown in Fig. 4.

The unit of viscosity is centipoise (cp) defined as (Skovborg et al. 1966).

Shear stress relates to the frictional forces within a fluid during flow. Wells and Merrill (1961) have estimated that the rate of shear in the aorta is 230.00 sec⁻¹ and in small arterioles and venules 11.50 sec⁻¹.

At the low shear rates, which exist under certain physiological conditions, hyperviscosity could result in reduced blood flow in the microcirculation even to the extent of complete blood stasis as shown in Fig. 5 and Table 1.

Clinical Manifestations

The infant usually appears plethoric, cyanotic, or both. The colour is somewhat more purplish than the usual cyanosis.

Cerebral signs include excessive irritability, stiffness, jitteriness, generalized or focal seizures or, conversely, severe lethargy, inability to suck, and marked hypotonia, associated at times with apnea and cyanosis (Wood, 1959; Kontras, 1972; Mackintosh and Walker, 1973; Gross et al., 1973).

Cardiopulmonary signs and symptoms like tachypnea, dyspnea with cardiac decompensation or spells of apnea (Naeye, 1963; Danks and Stevens, 1964; Gatti et al, 1966; O'Connor et al., 1968; Kontras, 1972; Gross et al., 1973; Fouron and Herbert, 1973). These signs and symp-
FIG. 4: Viscosity-Hematocrit curves at different shear rates (Adapted from Wells & Merriam, 1961).
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Symptoms may lead one to establish an erroneous diagnosis of cyanotic congenital heart disease. It is of additional interest and importance, however, that in Gatti's experience patients under the age of 1 month with cyanotic heart disease rarely have hematocrit values over 65%.

Fouron and Hebert (1973) believed that plethora with signs of cardiac failure are associated with polycythemic hypervolemia whereas cyanosis with cerebral signs and symptoms are associated with polycythemic normovolemia.

Papageorgiou and Stern (1972) reported a case of a newborn infant with gangrene of the right index finger in which the only abnormal finding other than the gangrenous finger was the high hematocrit (78%). They explained it as caused by hypoxia resulting from interfered red cell passage and tissue perfusion.

Humbert et al., (1969) described 2 male infants with preapism in whom polycythemia was the only detectable cause.

Signs and symptoms of hypoglycemia, increased bilirubin and coagulation defects may be noted because of sludging of blood flow.

Gross et al., (1973) presented the following histogram of signs and symptoms associated with hyperviscosity in 18 neonates (Figure 6).

Actually, the majority of infants with abnormally high hematocrits do not develop clinical disease which is not readily explained from the reported studies. Gatti et al., (1966) found neither cyanosis nor cardiopulmonary distress in 25 infants with hematocrit values greater than 75%.

Humbert et al., (1969) stated that: "certain groups of polycythemic infants seem more prone to develop associated complications than others" and found small for gestational age infants and male infants to be such groups. Fahey et al., (1965) suggested host factors which determine the vulnerability of individuals to elevated viscosity.

Laboratory and other related examinations

In some patients there may be roentgenographic evidence of cardiac enlargement and pulmonary vascular congestion with or without pleural fluid (Gatti et al., 1966; O'Connor et al., 1968; Lees, 1970). A simple chest roentgenogram can help to differentiate between polycythemia with hypervolemia and polycythemia without hypervolemia. In presence of hypervolemia and cardiac failure, cardiomegaly is a constant finding (Fouron and Hebert, 1973). However, the cardiovascular changes are transient and subside concurrently with the abnormal hematologic findings (Gatti et al., 1966). O'Connor et al., (1968) postulated that a portion or all of the blood normally distributed in the placenta is transfused.
FIG. 5: Viscosity/Shear rate curve (Adapted from Wells & Merrill, 1961).

Viscosity/Shear rate curve for whole blood and plasma.
into the infant at some time in the perinatal period resulting in hypervolemia, pulmonary congestion, cardiomegaly due to congenital heart failure and pleural effusion. Possible micro-emboli or thrombosis in the pulmonary circulation causes pleural fluid, pulmonary vascular congestion and cardiomegaly.

An abnormal electrocardiogram may be found during the period of cardiorespiratory distress.

An abnormal electroencephalogram may be found with or without seizures (Gross et al., 1973).

Funduscopical findings may be that of papilloedema, retinal hemorrhage, dilatation and tortuosity of the retinal vessels or central retinal vein closures (Meadows, 1947; Chrispin and Darke, 1962; Rothstein, 1972).

Guest and Brown in 1957 (cited by Kontras, 1972) recorded a mean cord blood hematocrit of 52.3%. They found a hematocrit of 58.2% on day 1, 54.5% on day 3 and 54.9% at 1 week. Moe (1965) reported normal capillary blood hemoglobin content, hematocrit level and erythrocyte count to be 19.8 gm% (S.D. = 2.4), 65.9% (S.D. = 7.5) and 5.40 mill./mm$^3$ (S.D. = 0.65) respectively, from 2 to 6 days of age. Sommer and Kontras (1971), cited by Kontras (1972) studying venous blood found average hematocrits of 55% ± 6.5 for day 1, 51% ± 5.91 for day 2, and 51% ± 5.98 for day 3. The source of sampling is to be taken into consideration. The average erythrocyte count is 804,000 per cm$^3$ and the hemoglobin 2.85 gm% higher in the capillary samples (Wood, 1959) and this marked capillary venous hematocrit difference is observed during the first 5 days of life. Heel warming improves the capillary venous hematocrit correlation (Oh and Lind, 1966).

The hematocrit can be determined by making use of a microhematocrit centrifuge.

Viscosity of heparinized (7 units per ml) whole blood (1.2 ml sample) can be determined using a Wells-Brookfield microviscometer with circulating water bath at 37° C. This apparatus can measure the shear stress at 8 different shear rates (230, 115, 46, 23, 11.5, 5.75, 2.3 and 1.15 sec$^{-1}$). The shear stress developed in the blood can be read from a dial. The viscosity is calculated by dividing the measured shear stress by the corresponding shear rate (Skovborg et al., 1966).

Filterability is a measure of erythrocyte deformability. Filterability of washed cells is measured as flow velocity (microlitres per second) during constant flow.

Considering the afore-mentioned pathophysiology, determination of blood unconjugated bilirubin, blood smear for signs of hemolysis, thrombocyte count and other coagulation studies pertinent to intravascular
### TABLE 1: Viscosity in the newborn (cp) (Mackintosh and Walker, 1973)

<table>
<thead>
<tr>
<th>Shear rate (sec⁻¹)</th>
<th>Mean 1 SD</th>
<th>1 SD</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.16</td>
<td>33.7</td>
<td>9.5</td>
</tr>
<tr>
<td>2.32</td>
<td>23.8</td>
<td>5.5</td>
</tr>
<tr>
<td>5.70</td>
<td>15.0</td>
<td>3.2</td>
</tr>
<tr>
<td>11.50</td>
<td>11.2</td>
<td>2.1</td>
</tr>
<tr>
<td>23.00</td>
<td>9.0</td>
<td>1.0</td>
</tr>
<tr>
<td>46.00</td>
<td>7.3</td>
<td>1.3</td>
</tr>
<tr>
<td>116.00</td>
<td>6.0</td>
<td>1.0</td>
</tr>
<tr>
<td>232.00</td>
<td>5.5</td>
<td>1.1</td>
</tr>
</tbody>
</table>

![FIG 6: Histogram of signs and symptoms associated with hyperviscosity in 18 neonates.](image_url)
coagulation should be done. Blood sugar, calcium and magnesium estimations should be done in the routine testing of the infant with cerebral signs.

To determine the cause of polycythemia, estimation of fetal hemoglobin, Beta₂A-globulin and differential red cell agglutinations are mandatory.

Prognosis

Most of the symptomatic cases reported in the literature improved dramatically after appropriate therapy. A lot of other cases were reported even with symptoms subsiding spontaneously (Gatti et al., 1966; Mackintosh and Walker, 1973). A long-term evaluation, however, of the symptomatic and asymptomatic cases is still necessary in view of subsequent neurologic impairment. It is noteworthy that insufficiency of cerebral perfusion may occur even in the absence of symptoms (Kontras, 1972). Cases with residual neurologic deficits reported in the literature are still too few. Of Wood's 2 patients who had convulsions associated with cyanosis and plethora, one developed grand mal seizures with episodes of transient blindness 7 years later (Gatti et al., 1966). Of the 3 patients in Gatti's 10 symptomatic study cases with myoclonic seizures, 2 had neurologic sequelae viz hemiparesis and myoclonic seizures at follow up. Gross et al. (1973) found motor and/or mental retardation on subsequent examination in 4 of his 10 symptomatic cases. Danks and Stevens (1964) found a slight internal strabismus developed in his patient at a later date.

Treatment

There is little agreement as to indications for treatment. Usually no treatment is necessary (Wood, 1959) and spontaneous improvement generally occurs (Gatti et al., 1966) but if there are symptoms of lethargy, anorexia, respiratory distress or convulsions, then treatment is indicated (Wood, 1959; Danks and Stevens, 1964; Kontras, 1972).

Hyperviscosity of blood is multifactorial and effective clinical management of the many circulatory and hematologic conditions represented by these syndromes will be based upon an understanding of mechanisms that create the hyperviscous state (Wells, 1970).

For a rational therapeutic approach it is important to know whether or not the polycythemic neonate is hypervolemic.

In normovolemic polycythemia there is a decreased cardiac output and partial exchange transfusion using 30 ml/kg body weight plasma or plasma expander may produce gratifying results when the venous hematocrit is 65 to 70% (Mackintosh and Walker, 1973). The capillary hematocrit is usually 5-6% higher. Exchange transfusion with plasma
or plasma expander has the advantage over exchange transfusion with saline or glucose saline because it eliminates the risk of hypovolemia.

In hypervolemic polycythemia there is an increased cardiac output and those infants suffering from cardiac failure may benefit from a phlebotomy of 10 per cent blood volume (Kontras, 1972) to bring the blood volume to a nearly normal value. Phlebotomy could be fatal to a cyanotic baby who has normovolemic polycythemia by decreasing an already very low cardiac output (Fouron and Hebert, 1973).

Administration of oxygen serves to reduce pulmonary vascular resistance (Gersony, 1973) and decreases the red cell volume and hematocrit value by depressing bone marrow activity thereby lowering the viscosity of the blood (Gatti et al., 1966).

Anticonvulsant drugs in presence of neurologic seizures and digitalis in presence of cardiac failure may be considered with regard to therapy.

Summary and conclusions

Syndrome of hyperviscosity in the neonate is a well-established condition. Polycythemia is defined as a venous hematocrit of 65% or greater. Polycythemic hyperviscosity is the most common and may be due to intrauterine anoxia, small for gestational age and dysmaturity, chromosomal abnormalities, maternofetal and twin-to-twin transfusions, and delayed cord clamping. Because of a variety of reasons, the newborn is particularly susceptible to developing hyperviscosity. Clinical manifestations are not always present and are pertinent to circulatory overload or sludging of blood flow. Early diagnosis and prompt treatment are highly desirable as the condition is life-threatening or may bring about sequelae whereas treatment is promising. A distinction between normovolemic and hypervolemic polycythemia should be attempted in regard to treatment.

REFERENCES


