A Vasoactive Inotropic Score Predicts the Severity of Hypotensive Shock and Mortality in Preterm Infants

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Research Article

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Abstract

Objective: To validate the prediction of the severity of hypotensive shock and mortality using the vasoactive inotropic score in preterm infants.

Methods: In this retrospective study we calculated the vasoactive inotropic score (VIS) and cumulative exposure to cardiovascular medications over time (VISct) in a cohort of preterm infants with hypotensive shock who received a cardiovascular support. Receiver operator curve was constructed to predict the primary outcome which was death due to hypotensive shock.

Results: VIS had an area under the curve of 0.73 (95% CI 0.85-0.98, \( p < 0.001 \)). A VIS cut off of 25 has sensitivity and specificity of 66% and 92%, and positive and negative predictive values of 78.5% and 83%, respectively.

Conclusion: High VIS predicts high mortality rate due to irreversible shock in preterm infants

What Is Known

Hypotensive shock severity increases mortality in premature infants

What is new: VIS predicts the mortality in preterm infants with hypotensive shock

Introduction

Management of hypotension is an unresolved challenge to neonatal intensive care practitioners(1). The concern about hypotension in preterm infants and the main rationale of neonatologists to treat, is the detrimental effect on end-organ perfusion(2). Hypotension has been associated with several adverse effects that reflect the cardiovascular instability and end-organ injury; the most common detrimental short term clinical consequences are intraventricular hemorrhage (IVH) and periventricular leukomalacia, necrotizing enterocolitis (NEC), and finally death (3). Between 16 to 98 % of preterm infants receive cardiovascular medications during admission(4).

Preterm infants who are both hypotensive and in shock - defined as low blood pressure with either lactic acidosis or oliguria - will probably have a high mortality. We hypothesized that a preterm infant with hypotension unresponsive to a certain dose of a cardiovascular drug as represented by the vasoactive inotropic score (VIS) can represent a marker at which the treating team should reassess the management strategy rather than continuing to increase the cardiovascular drug dosage or adding more drugs in the absence of clinical improvement. Blood pressure has never been proven to be a sensitive marker reflective of end organ blood flow especially when relying solely on mean blood pressure (MABP) and ignoring the physiologic values of systolic and diastolic blood pressures(5, 6). VIS may be useful as an independent predictor of poor short-term outcome after cardiothoracic surgery (7, 8) and predicts eventual morbidity and mortality in post-operative congenital heart surgery with cardiopulmonary bypass(9). The aim of this study was to validate the value of VIS for prediction of the severity of hypotensive shock and mortality in preterm
infants < 33 weeks. We hypothesized that a higher VIS score would be associated with high mortality in preterm infants < 33 weeks gestation.

**Methods**

All infants who were admitted to the 2 tertiary NICUs in Winnipeg, Canada between 2011–2015 who received cardiovascular medications were identified using pharmacy’s electronic records. Preterm infants < 33 weeks were identified and a retrospective chart review was undertaken. Eligibility criteria was premature babies < 33 weeks who had Clinical evidence of compromised systemic circulation with at least one of the following: low mean arterial blood pressure (MABP) as defined by less than the lower 95% confidence interval (CI) for the corrected gestational age as the standard monitoring, oliguria less than 1 mL/kg/h for at least 12 hours, or lactic acidosis greater than 2.8 mmol/l, and received cardiovascular medication (1, 10). Infants with congenital or chromosomal anomalies were excluded. In this study we calculated the (VIS) as follows [Dopamine (mcg/kg/min) + Dobutamine (mcg/kg/min) + Epinephrine (mcg/kg/min) ×100 + Norepinephrine (mcg/kg/min)×100 + Vasopressin (IU/kg/min)×10000 + Milrinone (mcg/kg/min)×10] (7, 9). We considered the highest dose for each drug if continued for > 6 hours. We calculated the cumulative vasoactive inotropic (VISct) exposure to the cardiovascular medications by multiplying the (VIS) of each drug with the exposure time in hours.

Demographic and clinical data were abstracted from the chart review. Those data included antenatal steroids, gestational age (GA), birth weight, sex, Apgar scores at 5 and 10 minutes, confirmed sepsis, and IVH. The cumulative time of compromised systemic circulation (CSC) during NICU admission was considered as marker of disease severity. CSC was considered present if the infant had one or more of the following: hypotension as described above, oliguria < 1ml/kg/hour for at least 12 hours beyond the first postnatal 12 hours, or lactic acid > 2.8mmol/dl(5, 6). The study period was before the targeted neonatal echocardiography was introduced in our units as a standard of care for assessment of infants with refractory shock; the cardiovascular medications were prescribed empirically starting with dopamine with the addition of dobutamine or epinephrine as second or third line, plus the administration of multiple empirical normal saline boluses. Our primary outcome was death due to hypotensive shock.

Ethics approval was obtained from University of Manitoba Research Ethics Board.

**Statistical analysis:**

SPSS v 24 (SPSS Inc, Chicago, Illinois) was used to perform the statistical analysis. Data are presented as median with interquartile range or frequencies. Comparisons between groups were analyzed by Mann-Whitney U test; Frequencies were analyzed using Chi-square; \( p < 0.05 \) was considered significant. The primary outcome for infants with hypotensive shock was evaluated using Cox proportional hazards regression model. Multivariate and adjusted multivariable analysis between groups was also performed using the general linear model to test effects between subjects and analyzed by multi-way ANOVA with Tukey's multiple comparisons test; \( p \) values < 0.05 were considered significant. A receiver operating characteristic (ROC) curve was constructed, and predictive values were calculated. Pearson correlation was
used to correlate \( \text{VIS} \geq 25 \) associated with death with other variables, with relevant graphs created by GraphPad program.

**Results**

One hundred and seventeen preterm infants < 33 weeks GA who had significant hypotension were enrolled in the study; 38 (32.5%) died and 79 (67.5%) survived. Table 1 showed comparison of the descriptive data between both groups. The groups had similar birth weights and gestational ages. Nine infants presented initially with lactic acidosis, 4 with oliguria and 8 with both, and 96 infants presented initially with hypotension. Infants who died have developed shock at an earlier postnatal age day 2 (1–8) vs. day 4 (1–14) \( (p < 0.05) \), and had a significantly higher VIS (median, IQR): 28 (12, 37) vs. 10(7, 18) \( (p < 0.0001) \) compared to those who recovered. Infant who died also had a higher fraction of inspired oxygen (FIO\(_2\)) at the time of the highest recorded VIS 0.45(0.36–0.65) vs. 0.4(0.3–0.5) \( (p < 0.05) \) and lower serum cortisol level 294(196–739) vs. 1640(383–1750) \( (p < 0.05) \). VIS value had an area under the curve of 0.73 (95% CI 0.85–0.98, \( p < 0.001 \)). A VIS value cut off of \( \geq 25 \) has sensitivity and specificity of 66% and 92%, and positive and negative predictive values of 78.5% and 83%, respectively, Fig. 1 shows the receiver operator curve.
Table 1
Descriptive data of the studied groups presented as median (IQR) and percentages, p-value is bold when significant

|                        | Hypotension in survivals | Hypotensive shock with death N = 38(32.5%) | p value |
|------------------------|--------------------------|---------------------------------------------|---------|
| **GA (wk)**            | 26 (25–29)               | 25(24–27)                                   | 0.07    |
| **Birth Weight (g)**   | 860(740–1200)            | 856(766–1056)                               | 0.2     |
| **Male Sex**           | 45(57%)                  | 26(68%)                                     | 0.5     |
| **Antenatal steroids** | 51(64%)                  | 23(60%)                                     | 0.6     |
| **Age at presentation (days)** | 4(1–14)             | 2(1–8)                                      | <0.05   |
| **Systolic blood pressure (mmHg)** | 33(29–37)     | 34(30–38)                                   | 0.3     |
| **Mean blood pressure (mmHg)** | 25(22–27)         | 24(21–25)                                   | 0.07    |
| **Diastolic blood pressure (mmHg)** | 18(17–22)        | 17(15–19)                                   | 0.12    |
| **Urine output hours (ml/kg/hr)** | 2.6(1.3–4.2) | 3(1.7-4)                                    | 0.1     |
| **Lactic acid hours (mmol/l)** | 1.9(1.2–3.4)     | 2.6(1.2–5.1)                                | 0.7     |
| **Confirmed Infection** | 26(32%)                  | 13(34%)                                     | 0.6     |
| **Vasoactive inotropic score (Highest)** | 10(7–18)          | 28(12–37)                                   | <0.0001|
| **Cumulative exposure to CVS medications VISct** | 312(116–912)   | 388(172–2088)                               | <0.01   |
| **Cumulative time of CSC (hours)** | 72(24–120)     | 62(24–121)                                  | 0.9     |
| **Serum cortisol level** | 1640(383–1750) | 294(196–739)                                | <0.05   |
| **Fraction of inspired oxygen** | 0.4(0.3–0.5) | 0.45(0.36–0.65)                            | <0.05   |
| **IVH (any grade)**    | 47 (59%)                 | 24 (63%)                                    | 0.3     |

A multivariate analysis was conducted and shows significant difference between survivors and infants who died before discharge on variables GA, VIS, hours of CSC, confirmed infection and VISct with Wilk’s Lambda = 0.75, F (4-112) = 9.3 p = 0.0001 partial $\eta^2$ =0.25. A separate ANOVA was conducted for each dependent variable, with each ANOVA evaluated at an alpha level of 0.05, both VIS and VISct were significantly higher in the non survivors ($M$ of VISct = 1827), ($M$ of VIS = 46) compared to survivors ($M$ of VISct = 850), ($M$ of VIS = 12), $p<0.05$, partial $\eta^2$=0.04 for VISct and $p<0.0001$, partial $\eta^2$=0.18 for VIS. There was no significant difference in GA ($p = 0.13$, partial $\eta^2$=0.02), there was no significant difference in hours with CSC between survival and non survival group ($p = 0.9$, partial $\eta^2$=0.0001). Figure 2 shows Cox proportional of death of infants with VIS ≥ 25 compared to infants with VIS < 25 and in relation to exposure time to cardiovascul
drugs. Table 2 and Fig. 3 show the association between infants with VIS who either dead or survived with the following variables; lactic acid, urine output, systolic blood pressure, diastolic blood pressure, mean blood pressures, serum cortisol levels, gestational age, and Apgar scores, there was significant correlation with lactic acid, urine output, systolic blood pressure, gestational age, and serum cortisol level. Higher percent of cases underwent delayed cord clamping was associated with VIS < 25, which was not the case with the percent of cases with confirmed infection Fig. 4.

Table 2
Correlation between duration of VIS ≥ 25 in infants who dead, and physiologic parameters:

| parameters                | CORRELATION |   |
|---------------------------|-------------|---|
|                           |             |   |
| Lactic acid               | 0.68        | 0.008** |
| Urine output              | 0.64        | 0.041*  |
| SYSTOLIC BLOOD PRESSURE   | 0.59        | 0.043*  |
| DIASTOLIC BLOOD PRESSURE  | 0.79        | 0.022*  |
| MEAN BLOOD PRESSURE       | 0.54        | 0.068   |
| SERUM CORTISOL LEVEL      | 0.71        | 0.032*  |

Discussion

We have examined the predictive value of the vasoactive inotropic score in preterm infants with hypotensive shock aiming to predict irreversible shock early which might alert the clinical team to consider different management strategy or consider to formulate a physiologic based medical recommendation by utilizing the new emerging techniques like targeted neonatal echocardiography and near infrared spectroscopy as an alternative to empiric approach(5, 6, 11). This study was conducted before implementing the physiologic based integrated hemodynamic approach, which might explain the high VIS in significant number of cases(5). The formulated medical recommendation could be either discontinuation of a cardiovascular medication causing side effects more than supporting systemic circulation, or consider more investigations to elucidate the pathophysiologic mechanism(12, 13). We found that VIS ≥ 25 was associated with higher mortality in preterm infants with hypotensive shock and is an alert number to start considering other modalities of treatment based on further imaging (TNE) Targeted Neonatal Echocardiography to assess underlaying pathophysiologic mechanism(14, 15). This is the first study to our knowledge describing the value of VIS in this population but it has been validated in older children with shock(8).

Current research confirms that the measurement of blood pressure does not reflect either tissue perfusion or oxygen delivery(16, 17), and current therapeutic interventions for hypotension are not associated with improved in-hospital outcomes or neurodevelopmental outcomes (18, 19). This could be related to lack of insight regarding the significance of physiologic markers which reflect variations in blood flow, together with
adopting empirical cardiovascular support without assessment of the underlying pathophysiology (12, 20). Identifying whether CSC a result of impaired preload, afterload, contractility, or the oxygen-carrying capacity (or a combination of those factors) can facilitate a more targeted and more meaningful approach to management, we analysed the preterm infants with CSC before implementing our hemodynamics protocol, which could explain high VIS in significant number of cases (21–23). We reported the impact of integrated evaluation of hemodynamics on clinical outcomes including VIS, and the time to clinical recovery and both were significantly lower in the group managed with the formulated pathophysiologic medical recommendation compared to infants managed with empiric approach (5). It is not uncommon that in refractory hypotensive shock, the number and/or dose of cardiovascular medications is empirically increased. this approach may overlook the detrimental side effect of these drugs, which may increase the risk of death in some situations (5, 6). VIS has been validated in neonates post cardiothoracic surgery and it was predictive of poor short-term outcomes (7, 24). The strengths of this study are using the VIS score as a marker of impending and irreversible shock and validating VISct which reflects the impact of the cumulative doses of cardiovascular drugs used before death or discharge. The limitations of this study are small sample size and the retrospective design and the lack of comparison with similar cases managed with physiologic approach.

**Conclusions**

VIS maybe useful in predicting severity of the hypotensive shock and mortality in preterm infants, we found that VIS \( \geq 25 \) was associated with higher mortality in preterm infants. This is the first study to our knowledge describing the value of VIS in this population.

**Abbreviations**

CSC: Compromised systemic circulation

IVH: Intraventricular hemorrhage

MABP: Mean arterial blood pressure

NEC: Necrotizing enterocolitis

VIS: Vasoactive inotropic score

VISct: Vasoactive inotropic score with cumulative time of exposure to cardiovascular medication

**Declarations**

**Funding:** No Funding was secured for this study.

**Conflicts of interest:** The authors have no financial relationships to disclose and no conflict of interest.

**Ethics approval:** was obtained from University of Manitoba Research Ethics Board
Consent to participate: (It was a retrospective chart review study)

Consent for publication: Obtained, The material is original research, has not been previously published and has not been submitted for publication elsewhere while under consideration.

Availability of data and materials: Available

Code availability: NA

Authors’ contribution:

Reem Amer; designed the study, collected the data, wrote the manuscript

Mary Seshia; reviewed the manuscript

Yasser Elsayed; Supervised the protocol design, reviews the statistical analysis, reviewed the manuscript.

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Figures
Diagonal segments are produced by ties.

Figure 1

ROC curve of VIS
Figure 2

Cox proportional of death of infants with VIS ≥ 25 compared to infants with VIS < 25 and in relation to exposure time to cardiovascular drugs
Figure 3

The association between infants with VIS who either died or survived with the following variables: lactic acid, urine output, systolic blood pressure, diastolic blood pressure, mean blood pressures, serum cortisol levels, gestational age, and Apgar scores, *= p value < 0.05, **= p value < 0.01.
Figure 4

The association between VIS and percent of cases with delayed cord clamping, and conformed infection, **
= p value < 0.01