Cardiac asystole at birth: Is hypovolemic shock the cause? *

J. Mercer.a,b,*, D. Erickson-Owensa,b, R. Skovgaardc

a College of Nursing, University of Rhode Island, 2 Heathman Road, Kingston, RI 02881, United States
b Adjunct, Brown University, United States
c Department of Obstetrics and Gynecology, University of Rochester at Highland Hospital, United States

SUMMARY

A birth involving shoulder dystocia can rapidly deteriorate—from a fetus with a reassuring tracing in the minutes before birth, to a neonate needing aggressive resuscitation. Infants experiencing a traumatic birth involving shoulder dystocia may be severely compromised, even when the preceding labor was uncomplicated. This paper presents two cases in which infants had normal heart beats recorded 5–10 min before birth and were born with cardiac asystole following shoulder dystocia. Often, in cases of shoulder dystocia, infants shift blood to the placenta due to the tight compressive squeeze of the body in the birth canal (along with cord compression) and thereby may be born hypovolemic. Our hypothesis is that the occurrence of sudden cardiac asystole at birth is due to extreme hypovolemic shock secondary to the loss of blood. At birth, the sudden release of pressure on the infant’s body results in hypoperfusion resulting in low central circulation and blood pressure. Severe hypovolemic shock from these effects leads to sudden cardiac arrest. Immediate cord clamping maintains the hypovolemic state by preventing the physiologic and readily available placental blood from returning to the infant. Loss of this blood initiates an inflammatory response leading to seizures, hypoxic-ischemic encephalopathy, and brain damage or death. Animal studies have shown that human umbilical stem cells injected into a rat’s abdomen after induced brain damage, can protect the rat’s brain from developing permanent injury. To prevent damage to newborns, the infant must receive the blood volume and stem cells lost at the time of descent and immediate cord clamping. Recommended countermeasures for research include: (1) resuscitation at the perineum with intact cord; or (2) milking the cord before clamping; or (3) autologous transfusion of placenta blood after the birth; or (4) rapid transfusion of O negative blood after birth and before seizures begin.

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sudden release of birth canal pressures, can result in sudden cardiaco arrest at birth. The authors use two cases involving shoulder dystocia and cardiac asystole to frame the discussion. Evidence to support this theory is reviewed and recommendations are provided for future research and immediate neonatal management in different settings.

**The cases**

The authors had access to birth records for these two cases as either an expert witness or from information shared by a defendant. Table 1 compares details of the two cases and highlights several points for discussion of the potential effect of shoulder dystocia on the neonate.

**Case 1: severe shoulder dystocia, seizures, and death**

A 35 year old G2 P1 with spontaneous rupture of membranes at 37\(\frac{1}{7}\) weeks was induced. The fetus appeared large upon admission. Her first birth was a vaginal delivery of a 3629 g male at term.

**Labor**

The first stage of labor was 6 h followed by a 2 h second stage. Pushing was compromised by a dense epidural. Vacuum assistance was used to gently pull the head through four contractions (the fetus had profuse hair and the cup placement did not achieve optimum suction). The external fetal monitor tracing throughout labor was essentially normal with the baseline rate 150–160 bpm and good accelerations. In the last hour, the baseline rate increased to 165–175 bpm.

**Delivery**

As the head began to deliver slowly, the fetus was observed by the physician, the mother, and the nurse to be making vigorous efforts to move its head. Over the next 30 s, the head was fully delivered. External fetal monitoring gear was removed with the last observed heart rate of 146–162 bpm immediately after the delivery of the head. The usual shoulder dystocia maneuvers were initiated. With significant effort the baby was successfully rotated using Wood’s screw maneuver and the posterior arm was delivered. Traction was required by two providers to deliver the chest and abdomen. The birth of the body occurred 6 min after the monitor was unhooked from the monitor. The cord was clamped and cut immediately and the pale, limp baby was passed to the resuscitation team.

**Resuscitation**

The baby weighed 4876 g and was born with Apgar scores of 0\(^1\), 0\(^5\), 0\(^10\). Efforts to resuscitate the infant included a full cardio-pulmonary resuscitation. Endotracheal intubation was performed by the anesthesiologist at 1 min of age. Endotracheal epinephrine was given on four occasions: at 1, 6, 14 and 18 min. Within 3 min of the birth, a pediatrician was present. After 15 min, the baby was moved to the nursery and an umbilical venous (UV) catheter was placed. At 16 min 50 mL of normal saline was given with 8 mL of bicarbonate solution. At 18 min epinephrine 0.3 mg was given by IV line. An additional 4 mL of bicarbonate was given at 22 min. At 23 min, a heart rate (HR) was observed. The first arterial blood gas (ABG) was collected from the radial artery at 31 min of life, showing pH 6.73, pCO2 112 PO2 of 65 and O2 saturation of 66%. Blood pressure (BP) was 69/39 with a HR of 160, but at 1.5 h the BP dropped to 51/21 prompting the initiation of dopamine at 5 mcg/kg/minute. At 1 h of age seizure activity was noted and phenobarbital therapy initiated.

**NICU Course**

The infant was transferred to the Level III Neonatal Critical Care Unit of a nearby hospital. Admission vital signs on dopamine included HR 121, BP 55/24, and temperature of 37.5 °C. The baby was intubated and on a ventilator. The first hematocrit and hemoglobin (H&H) noted at five hours of age was 43.7% and 14.7 g/dL. Vital functions of respiration, renal function, temperature control, and BP stability recovered by three days of age. Dopamine was discontinued. There was profound brain damage.

**Outcome**

The infant had severe feeding problems with poor suck and frequent aspiration necessitating a PEG tube. Regurgitation and aspiration of food remained a major problem. At three weeks of age the child had not recovered any significant neurologic function. He was discharged home with hospice support. Death at home at two months of age was secondary to aspiration pneumonia. No placental pathology was noted. An autopsy was not performed.

**Case 2: severe shoulder dystocia, seizures and cerebral palsy**

A 22 year old G1P0 at 41\(\frac{4}{7}\) weeks was induced after a normal prenatal course.

**Labor**

First stage was progressive over 11 h. Second stage lasted 1 h. The fetal heart tracing was reactive throughout labor with some tachycardia (160–180 bpm) and marked variability in the last half hour.

**Delivery**

The fetal head was slow to emerge. Time from delivery of the head to the delivery of the body was estimated to be between 5–10 min. A nuchal cord was present and was cut about two minutes before delivery.

**Resuscitation**

The infant appeared floppy, pale and was unresponsive. Apgar scores were 0\(^1\), 0\(^5\), 0\(^10\). The infant was bagged and masked and then intubated within three minutes. Pediatrics was in attendance by 15 min. The infant weighed 4214 g. A cord blood gas showed pH 7.11, CO2 67, base deficit of –9.6. Two doses of epinephrine were given and a heart rate of 100 was heard after the second dose at 18 min after birth. The infant was taken to the nursery and placed on a ventilator. On admission to the nursery, an ABG revealed a pH of 6.9. An umbilical artery catheter was placed and albumin was given twice. The systolic BP fell to 40 mHg and dopamine was started. Generalized seizure activity (bicycling and lip smacking) was noted between one and two hours. The infant was transferred

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**Table 1**

Commonalities between two cases of hypoxic ischemic encephalopathy.

<table>
<thead>
<tr>
<th></th>
<th>Case 1</th>
<th>Case 2</th>
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</thead>
<tbody>
<tr>
<td>Length of labor</td>
<td>6</td>
<td>8</td>
</tr>
<tr>
<td>1st stage</td>
<td>2</td>
<td>1</td>
</tr>
<tr>
<td>2nd stage</td>
<td>146–162 bpm</td>
<td>160–180 bpm</td>
</tr>
<tr>
<td>Last record heart rate</td>
<td>10 lbs 12 oz</td>
<td>9 lbs 4 oz</td>
</tr>
<tr>
<td>Infant’s birth weight</td>
<td>6 min</td>
<td>5–10 min</td>
</tr>
<tr>
<td>Time from birth of head to delivery of body</td>
<td>Immediate</td>
<td>Before delivery of shoulders</td>
</tr>
<tr>
<td>Cord clamping</td>
<td>6.9 @ 30 min</td>
<td>7.11 (cord) 6.72 @ 60 min</td>
</tr>
<tr>
<td>Cord or first pH</td>
<td>23 min</td>
<td>18 min</td>
</tr>
<tr>
<td>Length of asystole</td>
<td>1 h</td>
<td>1.5 h</td>
</tr>
<tr>
<td>First seizure</td>
<td>Severe</td>
<td>Severe</td>
</tr>
<tr>
<td>Hypotension</td>
<td>47% @ 5 h</td>
<td>38% at 1 h</td>
</tr>
</tbody>
</table>
to a Level III Nursery. Initial H&H was 38% and 13 g/dL. Examination of the placenta revealed no pathology.

Outcome
The child survived with CP (can walk but not run) and developmental delay. The child attends school but is three years below grade level. Experts offered that there was clear evidence of acute perinatal damage on MRI [5].

The hypothesis
Our hypothesis is that the sudden cardiac asystole noted in both of these cases is due to severe hypovolemic shock. The mechanism involved is a compressive squeeze of the body and umbilical cord as the large infant descends through the birth canal. This compression causes some volume of blood to be retained in the placenta rather than returning to the infant. At birth, when the baby's body is freed, there is a sudden release of the pressure especially on the infant's peripheral circulation. With insufficient blood supply to perfuse the entire body and maintain the central circulation, the heart stops beating. The usual practice of immediate cord clamping prevents the placental circulation from restoring the lost volume in a homeostatic fashion. The result is a severely compromised infant, who if resuscitated, may experience hypoxic-ischemic encephalopathy (HIE) as a result of low blood volume, the loss of red blood cells (RBCs) for oxygenation and hematopoietic stem cells essential for healing and repair.

We have been unable to find any current hypotheses to explain the mechanism of sudden cardiac asystole after birth in the medical literature. The resulting damage, hypoxic-ischemic cerebral injury, is believed to be a result of impaired cerebral blood flow occurring as a consequence of "interruption in placenta blood flow and gas exchange" with which we agree [6]. Most focus is given to what happens at the cellular level. Volpe discusses implications of reduced cerebral blood volume but does not link it to the infant's overall general blood volume [7]. The following discussion dissects each part of the hypothesis and provides evidence as available.

The second stage “squeeze”
Blood volume loss, significant enough to cause hypovolemic shock at birth, may occur as the infant descends into the birth canal prior to shoulder dystocia. The mechanism may be the compressive squeeze of the infant as it traverses the tight birth canal places pressure on the umbilical cord with or without a cord entanglement [8]. The soft-walled umbilical vein is more easily occluded than the arteries [9,10]. The muscular-walled, high-pressure arteries continue to move blood from the fetus to the placenta, while return flow from the placenta to the fetus in the thin-walled vein can easily be impeded. Thus there is a net transfer of blood volume from the fetus to the placenta. Few changes other than tachycardia may be seen on the fetal monitor tracing. Yet the stage is set for dramatic compromise at birth when the surrounding pressure is released and the neonate is left severely volume depleted.

Effects of vaginal wall pressure
Compensating for the loss of blood volume, pressure from the vaginal walls acts as an anti-shock garment to maintain blood pressure, circulation, and HR while the infant is in the vagina. The vaginal wall pressure most likely supports a transfer of blood from the peripheral circulation to the central circulation (brain, heart, lungs, kidneys) allowing the infant to maintain a near normal heart rate [11]. Anti-shock garments have been shown to provide 20–50 mmHg pressure to the lower body [12]. As the infant is born, there is sudden loss of the supporting vaginal wall pressure of the birth canal. This loss of surrounding pressure is akin to suddenly removing an anti-shock garment—it creates an immediate opening of the peripheral circulation and thus reduces the central circulation resulting in a drop of blood pressure (BP). The reduction in central circulation decreases blood flow to the heart causing hypoxia/anoxia and it immediately stops beating [11].

Hypovolemic shock
Hypovolemic shock due to inadequate circulating blood volume has the potential to cause profound damage quickly and consistently, including asystole. Hypovolemia, hypotension, hypovolemic shock and anemia have been documented in infants born with nuchal cords that were cut before delivery of the shoulders [8,13–15]. CP has been reported in cases when a nuchal cord is cut and then shoulder dystocia arises even with only a brief delay in the delivery [16–18]. Reports of reduction in birth weight with early clamping support the theory that this intervention can reduce the amount of placental transfusion [19–21].

Sudden cardiac arrest
The volume depleted infant is born in a state of hypovolemic shock and does not have enough blood in the central circulation to adequately oxygenate and perfuse the heart [22]. The heart suddenly stops beating secondary to hypoxia, resulting in immediate cardiac arrest [23]. Both of our case infants had normal heart rates recorded less than 6–10 min prior to the arrest. One infant was seen moving his head 6 min before birth yet was born lifeless.

Immediate cord clamping
Immediate cord clamping (ICC) leaves the infant with only the blood volume that was in his body at the moment of birth. ICC after a normal birth has been shown to result in a 30% reduction in blood volume and a 50% deficit of RBCs [24]. Blood loss or the blood diverted to the placenta due to the birth canal squeeze also creates a significant reduction of blood volume at birth.

Infants who undergo immediate cord clamping after a normal birth are left with about 70 mL/kg while infants with delayed cord clamping obtain about 90 mL/kg [24]. For a 4000 g infant, this difference is 280 mL versus 360 mL for the circulating blood volume [24]. Thus, delayed cord clamping or milking the cord will provide about 80 mL of blood for circulatory adjustments for an infant of this size. In the first few minutes after birth, the pulmonary capillary bed becomes a huge reservoir that must be rapidly perfused for the first time as the cardiac output to the lung shifts from 8% in utero to 50% as breathing is established [25]. This 80 mL of blood is not extra blood—it has been the respiratory blood in utero [26].

Loss of red blood cells. Yao and colleagues found that infants with a delay in cord clamping of three minutes obtain approximately 50% more RBCs [24]. RBCs play a critical role in oxygen and carbon dioxide transport. Jones estimated that infants need approximately 45 mg/kg RBCs for adequate oxygen-carrying capacity [27]. The only infants in the study of Yao and colleagues who obtained 45 mg/kg RBCs were those infants whose clamping was delayed for three minutes [24]. Whatever amount of blood is trapped in the placenta because of the second stage squeeze would add to RBC loss.

Measurement of blood volume and red cell volume. It is worth noting that accurate measurement of blood volume and red cell volume is notoriously difficult. The hematocrit level, being simple to obtain, is most commonly used. But it can be unreliable in circumstances of trauma or stress. Faxelius et al. measured red cell
volume by tagging the RBCs with non-radioactive chromium and found that infants with a history of asphyxia often had a low red cell volume even if no signs of blood loss were present [28]. Others studied blood volume in the 1st hour in asphyxiated infants using 131I tagged albumin and found larger blood volumes in asphyxiated babies [29]. However, Yao points out that the infants she studied had asphyxia livida (plethoric) and that infants with asphyxia pallida (pale) may have less blood volume. Better techniques to assess infant blood volume are needed to answer these questions. Measurement of superior vena cava flow may offer some insight but has only been reported in preterm infants [30].

Inflammation and stem cells

Two separate animal studies have demonstrated inflammation caused by blood loss alone [31,32]. Meier and colleagues (2006) demonstrated that loss of blood volume alone to the brain results in inflammation. Through ligation of the rat pup’s carotid artery, a progressive inflammatory process was created leading to neuronal death in the brain and resulting in spastic paresis. When human UCB cells were injected into the abdomens of affected rat pups within 24 h of injury, spastic paresis did not develop in the experimental group. Administering the UCB cells to the damaged pups later than 24 h did not prevent damage. Upon histologic examination of the rat brains at 21 days after the injury, Meier et al. [32] found that the human UCB stem cells crossed the blood-brain barrier, surrounded and infiltrated the damaged areas and appeared to provide scaffolding for repair. These cells were not found in any intact, uninjured areas suggesting that inflammatory factors (cytokines) from the damage areas signaled the stem cells [33]. We estimated that our case infants lost about one billion stem cells because of second stage compression compounded by immediate cord clamping. Might the stem cells have played a role in healing their deficits? [34].

Resuscitation

Developing evidence highlights perfusion as the key factor in adult resuscitation, such as in cardiac arrest. The 2005 American Heart Association guidelines for CPR places increasing importance on chest compressions over ventilations [35]. Ewy advocates re-conceptualizing CPR as “cardiocerebral” resuscitation (CCR) and recommends continuous chest compressions (CCC–CPR), without interruption for ventilations [36]. This recommendation is based on animal models in which cardiac arrest survival increased to 80% when CCC–CPR was applied, compared to 13% with standard CPR [37]. Maintenance of perfusion to the heart and brain was the critical factor in protecting function of those organs and impacting survival. Hypovolemic infants, such as our case infants, lack the crucial blood volume needed to adequately perfuse the heart and brain.

Research needed: Preventing hypovolemic shock at birth

The sight of the pale white, toneless infant at birth invokes fear in all caregivers. These infants are very difficult to resuscitate. The fear of managing them is real and appropriate. Often, a properly conducted resuscitation revives a terminally damaged infant (as in our first case) or does not revive the infant at all. Assisting in the delivery of the infant, holding the infant at the perineum for drying, stimulation, and resuscitation offers the opportunity for perfusely, there is no reason to suspect separation of the placenta.

Resuscitation at the perineum

Keeping a depressed infant at the perineum seems counter-intuitive. However, a full resuscitation can be done at the perineum without cutting the cord. Holding the infant at the perineum for drying, stimulation, and resuscitation offers the opportunity for some of the blood volume to return to the infant. This method allows the infant to get his own hemoglobin F, stem cells and the appropriate cytokine cocktail to guide the stem cells. The placental system continues to circulate in the first few minutes of life until the cord stops pulsating [41,42]. Unless the mother is bleeding profusely, there is no reason to suspect separation of the placenta.

Milking the cord

This option involves lowering the infant as much as possible and vigorously milking the cord two to four times before cutting it and taking the infant to the warmer [43,44]. Milking the umbilical cord may offer a viable option to prevent hypovolemic shock at birth when time and speed are critical factors. A review of the literature on umbilical cord milking or stripping supports this technique as a safe intervention resulting in higher neonatal hemoglobin levels and red cell volume at birth.

In nine controlled trials involving 803 babies (1949–2008), 350 infants had immediate cord clamping, 116 infants had delayed cord clamping, and 337 infants had umbilical cord milking [43–51]. Infants with immediate cord clamping had significantly lowered H&H levels than infants with delayed cord clamping or cord milking. No increase in jaundice or symptomatic polycythemia was reported among the infants in the studies.

Autologous transfusion of cord blood

The third intervention involves collecting cord blood under sterile conditions, using 60 cc syringes with appropriate additive, from the maternal side of the clamped umbilical cord. The blood can then immediately be transferred to the infant, at a volume of 15–20 mL/kg. The safety and efficacy of this intervention has been documented [52,53]. We believe that it will be imperative that the blood volume lost should be returned to the infant before one hour of age or about the time that seizures may begin (in our two cases seizures began at 60–90 min-see Table 1).
Transfusion of fresh whole blood

Given strong evidence that severe volume depletion can cause hypoxia resulting in serious and irreversible injury to newborns, a transfusion as soon as possible with fresh whole O negative blood at 10–20 mL/kg may be indicated when the previous measures are impossible or ineffective. Though this is the least optimal method, it may have helped our case infants and others like them as they had low hematocrit levels for term infants and needed vasoressors to maintain blood pressure. It does not provide the infant with hemoglobin F or activated stem cells, but it does supply red blood cells with oxygen carrying capacity and volume.

Caveat: cord blood gases

There is no immediate benefit to the infant for cutting the cord early in order to obtain blood gases. The infant is resuscitated by clinical appearance and responses while a result of any cord gases obtained is pending. We acknowledge that there may be some clinical advantage later in care but the irreversible damage may have occurred [54]. The act of cutting the cord to obtain the gases prevents any opportunity to correct the infant’s acid-base balance by extra-uterine resuscitation with intact cord. In addition, it limits transfusion of blood in cases where the infant is volume depleted, setting up a likelihood of additional hypoxic-ischemic injury of major organs. Further study may reveal that, when indicated, obtaining venous and arterial samples from an intact cord may prove safe and useful. Our case infants may have had different outcomes with resuscitation using their own blood.

Summary

In cases of shoulder dystocia, infants shift blood to the placenta due to the tight squeeze of the birth canal and thereby are born hypovolemic. At birth, the sudden release of pressure results in hypoperfusion and low blood pressure. Severe hypovolemic shock from these effects can result in sudden cardiac arrest. Immediate cord clamping maintains the hypovolemic state and delays treatment until volume expanders can be given artificially through an umbilical catheter instead of using physiologic and readily available placental blood. Blood loss sets off an inflammatory cascade which can result in seizures, HIE and later brain damage. To prevent these conditions, the infant must obtain the blood volume lost with the squeeze. Recommended countermeasures for research are: (1) resuscitation at the perineum with intact cord; or (2) milking the cord before clamping; or (3) autologous transfusion of placenta blood after the birth; or (4) rapid transfusion of O negative blood.

Acknowledgment

We are grateful to Dr. Peter Reagan who sent us a case description and raised the importance of publishing the case to open discussion of these vital issues. We dedicate this chapter to all the infants, like Nicholas, who lost their lives to teach us.

References